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91003, F-9277 Issy-les-Moulineaux (FR). LESAGE, Anne, Simone, Josephine [BE/BE]; Janssen Pharmaceutica N.V, Turnhoutseweg 30, B-2340 Beerse (BE).

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- (71) Applicant (for all designated States except US): JANSSEN PHARMACEUTICA N.V. [BE/BE]; Turnhoutseweg 30, B-2340 Beerse (BE).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): MABIRE, Dominique, Jean-Pierre [FR/FR]; Janssen-Cilag S.A., 1, rue Camille Desmoulins, TSA 91003, F-9277 Issy-les-Moulineaux (FR). VENET, Marc, Gaston [FR/FR]; Janssen-Cilag S.A., 1, rue Camille Desmoulins, TSA 91003, F-9277 Issy-les-Moulineaux (FR). COUPA, Sophie [FR/FR]; Janssen-Cilag S.A., 1, rue Camille Desmoulins, TSA 91003, F-9277 Issy-les-Moulineaux (FR). PONCELET, Alain, Philippe [FR/FR]; Janssen-Cilag S.A., 1, rue Camille Desmoulins, TSA

- (74) Agent: VAN BORM, Werner; Janssen Pharmaceutica N.V., Turnhoutseweg 30, B-2340 Beerse (BE).
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(54) Title: METABOTROPIC GLUTAMATE RECEPTOR ANTAGONISTS

(57) Abstract: The present invention concerns compounds of formula. In a preferable embodiment, X represents O; R1 represents C₁₋₆alkyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃-₁₂alkyl-moiety optionally may be replaced by C_{1-6} alkyloxy, aryl, halo or thienyl; R^2 represents hydrogen; halo; C_{1-6} alkyl or amino; R^3 and R^4 each independently represent hydrogen or $C_{1.6}$ alkyl; or R^2 and R^3 may be taken together to form $-R^2-R^3$, which represents a bivalent radical of formula -Z₄-CH₂-CH₂-CH₂- or -Z₄-CH₂-CH₂- with Z₄ being O or NR¹¹ wherein R¹¹ is C₁₋₆alkyl; and wherein each bivalent radical is optionally substituted with $C_{1.6}$ alkyl; or R^3 and R^4 may be taken together to form a bivalent radical of formula -CH₂-CH₂-CH₂-CH₂-; R⁵ represents hydrogen; Y represents O; and aryl represents phenyl optionally substituted with halo. The invention also relates to the use of a compound according to the invention as a medicament and in the manufacture of a medicament for treating or preventing glutamate-induced diseases of the central nervous system, as well as formulations comprising such a compound and processes for preparing such a compound.

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METABOTROPIC GLUTAMATE RECEPTOR ANTAGONISTS

The present invention is concerned with quinoline and quinolinone derivatives showing metabotropic glutamate receptor antagonistic activity and their preparation; it further relates to compositions comprising them, as well as their use as a medicine.

The neurotransmitter glutamate is considered to be the major excitatory neurotransmitter in the mammalian central nervous system. The binding of this neurotransmitter to metabotropic glutamate receptors (mGluRs), which are a subfamily of the G-protein-coupled receptors and which comprise 8 distinct subtypes of mGluRs, namely mGluR1 through mGluR8, activates a variety of intracellular second messenger systems. The mGluRs can be divided into 3 groups based on amino acid sequence homology, the second messenger system utilized by the receptors and the pharmacological characteristics. Group I mGluRs, which comprises mGluR subtype 1 and 5, couple to phospholipase C and their activation leads to intracellular calcium-ion mobilization. Group II mGluRs (mGluR2 and 3) and group III mGluRs (mGluR4, 6, 7 and 8) couple to adenyl cyclase and their activation causes a reduction in second messenger cAMP and as such a dampening of the neuronal activity. Treatment with Group I mGluR antagonists has been shown to translate in the presynapse into a reduced release of neurotransmitter glutamate and to decrease the glutamate-mediated neuronal excitation via postsynaptic mechanisms. Since a variety of pathophysiological processes and disease states affecting the central nervous system are thought to be due to excessive glutamate induced excitation of the central nervous system neurons, Group I mGluR antagonists could be therapeutically beneficial in the treatment of central nervous sytem diseases.

WO 99/26927 discloses antagonists of Group I mGlu receptors for treating neurological diseases and disorders, based – among others – on a quinoline structure.

WO 99/03822 discloses bicyclic metabotropic glutamate receptor ligands, none of them based on a quinoline or quinolinone structure.

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The present invention concerns compounds of formula

$$R^1$$
 R^4
 R^3
 R^2
 R^2
 R^3
 R^4
 R^3
 R^4
 R^3
 R^3
 R^5

an N-oxide form, a pharmaceutically acceptable addition salt, a quaternary amine and a stereochemically isomeric form thereof, wherein

5 X represents O; $C(R^6)_2$ with R^6 being hydrogen, aryl or C_{1-6} alkyl optionally substituted with amino or mono- or di(C_{1-6} alkyl)amino; S or N-R⁷ with R⁷ being amino or hydroxy;

R¹ represents C₁₋₆alkyl; aryl; thienyl; quinolinyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein the cycloC₃₋₁₂alkyl moiety optionally may contain a double bond and wherein one carbon atom in the cycloC₃₋₁₂alkyl moiety may be replaced by an oxygen atom or an NR⁸-moiety with R⁸ being hydrogen, benzyl or C₁₋₆alkyloxycarbonyl; wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyl, hydroxyC₁₋₆alkyl, haloC₁₋₆alkyl, aminoC₁₋₆alkyl, hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, halo, C₁₋₆alkyloxycarbonyl, aryl, amino, mono— or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, halo, piperazinyl, pyridinyl, morpholinyl, thienyl or a bivalent radical of formula –O-, -O-CH₂-O or — O-CH₂-CH₂-O-;

or a radical of formula (a-1)

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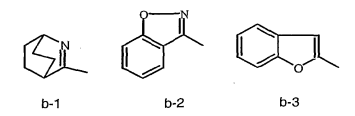
$$Z_1$$
 CH Z_2 (CH₂)_n

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wherein Z₁ is a single covalent bond, O, NH or CH₂;
Z₂ is a single covalent bond, O, NH or CH₂;
n is an integer of 0, 1, 2 or 3;
and wherein each hydrogen atom in the phenyl ring independently may optionally be replaced by halo, hydroxy, C₁₋₆alkyl,
C₁₋₆alkyloxy or hydroxyC₁₋₆alkyl;

or X and R¹ may be taken together with the carbon atom to which X and R¹ are attached to form a radical of formula (b-1), (b-2) or (b-3);

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R² represents hydrogen; halo; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkylthio; C₁₋₆alkylcarbonyl; C₁₋₆alkyloxycarbonyl; C₁₋₆alkylcarbonyloxyC₁₋₆alkyl; 5 C₂₋₆alkenyl; hydroxyC₂₋₆alkenyl; C₂₋₆alkynyl; hydroxyC₂₋₆alkynyl; tri(C₁₋₆alkyl)silaneC₂₋₆alkynyl; amino; mono- or di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxyC₁₋₆alkyl)amino; mono– or di(C₁₋₆alkylthioC₁₋₆alkyl)amino; aryl; $arylC_{1-6}alkyl; arylC_{2-6}alkynyl; C_{1-6}alkyloxyC_{1-6}alkylaminoC_{1-6}alkyl;$ aminocarbonyl optionally substituted with C₁₋₆alkyl, C₁₋₆alkyloxyC₁₋₆alkyl, 10 C_{1-6} alkyloxycarbonyl C_{1-6} alkyl or pyridinyl C_{1-6} alkyl; a heterocycle selected from thienyl, furanyl, pyrrolyl, thiazolyl, oxazolyl, imidazolyl, isothiazolyl, isoxazolyl, pyrazolyl, pyridyl, pyrazinyl, pyridazinyl, pyrimidinyl, piperidinyl and piperazinyl, optionally N-substituted with C₁₋₆alkyloxyC₁₋₆alkyl, morpholinyl, thiomorpholinyl, dioxanyl or dithianyl; 15 a radical –NH-C(=O)R⁹ wherein R⁹ represents

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 $C_{1\text{-}6}$ alkyloxycarbonyl, aryl, aryloxy, thienyl, pyridinyl, mono- or $di(C_{1\text{-}6}$ alkyl)amino, $C_{1\text{-}6}$ alkylthio, benzylthio, pyridinylthio or pyrimidinylthio; cyclo $C_{3\text{-}12}$ alkyl; cyclohexenyl; amino; arylcyclo $C_{3\text{-}12}$ alkylamino; mono- or $di(C_{1\text{-}6}$ alkyl)amino; mono- or $di(C_{1\text{-}6}$ alkyloxycarbonyl $C_{1\text{-}6}$ alkyl)amino; mono- or $di(C_{1\text{-}6}$ alkyloxycarbonyl)amino; mono- or $di(C_{2\text{-}6}$ alkenyl)amino; mono- or $di(C_{2\text{-}6}$ alkenyl)amino; mono- or $di(C_{2\text{-}6}$ alkenyl; piperididinyl; piperazinyl; indolyl; furyl; benzofuryl; tetrahydrofuryl; piperididinyl; pyridinyl; pyrazinyl; aryl; aryl $C_{1\text{-}6}$ alkylthio or a radical of formula (a-1);

 C_{1-6} alkyl optionally substituted with cyclo C_{3-12} alkyl, C_{1-6} alkyloxy,

a sulfonamid -NH-SO₂-R¹⁰ wherein R¹⁰ represents C_{1-6} alkyl, mono- or poly halo C_{1-6} alkyl, aryl C_{1-6} alkyl, aryl C_{2-6} alkenyl, aryl, quinolinyl, isoxazolyl or di(C_{1-6} alkyl)amino;

R³ and R⁴ each independently represent hydrogen; halo; hydroxy; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkyloxyC₁₋₆alkyl; C₁₋₆alkylcarbonyl; C₁₋₆alkyloxycarbonyl;

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 $C_{2\text{-}6}$ alkenyl; hydroxy $C_{2\text{-}6}$ alkenyl; $C_{2\text{-}6}$ alkynyl; hydroxy $C_{2\text{-}6}$ alkynyl; tri $(C_{1\text{-}6}$ alkyl)silane $C_{2\text{-}6}$ alkynyl; amino; mono— or di $(C_{1\text{-}6}$ alkyl)amino; mono— or di $(C_{1\text{-}6}$ alkyl)amino; mono— or di $(C_{1\text{-}6}$ alkyl)amino; aryl; morpholinyl $C_{1\text{-}6}$ alkyl or piperidinyl $C_{1\text{-}6}$ alkyl; or

- R² and R³ may be taken together to form -R²-R³-, which represents a bivalent radical of formula -(CH₂)₃-, -(CH₂)₄-, -(CH₂)₅-, -(CH₂)₆-, -CH=CH-CH=CH-,
 -Z₄-CH=CH-, -CH=CH-Z₄-, -Z₄-CH₂-CH₂-CH₂-, -CH₂-Z₄-CH₂-CH₂-,
 -CH₂-CH₂-Z₄-CH₂-,
 -CH₂-CH₂-Z₄-, -Z₄-CH₂-CH₂-, -CH₂-Z₄-CH₂- or -CH₂-CH₂-Z₄-, with Z₄ being
 O, S, SO₂ or NR¹¹ wherein R¹¹ is hydrogen, C₁₋₆alkyl, benzyl or C₁₋₆alkyloxycarbonyl; and wherein each bivalent radical is optionally substituted
 - or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH=CH-CH=CH- or -CH₂-CH₂-CH₂-CH₂-;
- 15 R⁵ represents hydrogen; cycloC₃₋₁₂alkyl; piperidinyl; oxo-thienyl; tetrahydrothienyl, arylC₁₋₆alkyl; C₁₋₆alkyloxyC₁₋₆alkyl; C₁₋₆alkyloxycarbonylC₁₋₆alkyl or C₁₋₆alkyl optionally substituted with a radical C(=O)NR_xR_y, in which R_x and R_y, each independently are hydrogen, cycloC₃₋₁₂alkyl, C₂₋₆alkynyl or C₁₋₆alkyl optionally substituted with cyano, C₁₋₆alkyloxy, C₁₋₆alkyloxycarbonyl, furanyl, pyrrolidinyl, benzylthio, pyridinyl, pyrrolyl or thienyl;
 - Y represents O or S:

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with C₁₋₆alkyl.

or Y and R⁵ may be taken together to form =Y-R⁵- which represents a radical of formula

-CH=N-N= (c-1);

-N=N-N= (c-2); or

-N-CH=CH- (c-3);

- aryl represents phenyl or naphthyl optionally substituted with one or more substituents selected from halo, hydroxy, C_{1-6} alkyl, C_{1-6} alkyloxy, phenyloxy, nitro, amino, thio, C_{1-6} alkylthio, halo C_{1-6} alkyl, polyhalo C_{1-6} alkyl, polyhalo C_{1-6} alkyloxy,
- hydroxyC₁₋₆alkyl, C₁₋₆alkyloxyC₁₋₆alkyl, aminoC₁₋₆alkyl, mono-or di(C₁₋₆alkyl)amino; mono-or di(C₁₋₆alkyl)aminoC₁₋₆alkyl, cyano, -CO-R¹², -CO-OR¹³, -NR¹³SO₂R¹², -SO₂-NR¹³R¹⁴, -NR¹³C(O)R¹², -C(O)NR¹³R¹⁴, -SOR¹², -SO₂R¹²; wherein each R¹², R¹³ and R¹⁴ independently represent C₁₋₆alkyl; cycloC₃₋₆alkyl; phenyl; phenyl substituted with halo, hydroxy, C₁₋₆alkyl,
- C₁₋₆alkyloxy, haloC₁₋₆alkyl, polyhaloC₁₋₆alkyl, furanyl, thienyl, pyrrolyl, imidazolyl, thiazolyl or oxazolyl;

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and when the R^1 -C(=X) moiety is linked to another position than the 7 or 8 position, then said 7 and 8 position may be substituted with R^{15} and R^{16} wherein either one or both of R^{15} and R^{16} represents C_{1-6} alkyl, C_{1-6} alkyloxy or R^{15} and R^{16} taken together may form a bivalent radical of formula -CH=CH-CH=CH-.

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As used in the foregoing definitions and hereinafter C_{1-6} alkyl as a group or part of a group encompasses the straight and branched chain saturated hydrocarbon radicals having from 1 to 6 carbon atoms such as, for example, methyl, ethyl, propyl, butyl, pentyl or hexyl; C_{2-6} alkenyl as a group or part of a group encompasses the straight and branched chain hydrocarbon radicals having from 2 to 6 carbon atoms and having a double bond such as ethenyl, propenyl, butenyl, pentenyl, hexenyl, 3-methylbutenyl and the like; C_{2-6} alkynyl as a group or part of a group defines straight or branched chain hydrocarbon radicals having from 2 to 6 carbon atoms and having a triple bond such as ethynyl, propynyl, butynyl, pentynyl, hexynyl, 3-methylbutynyl and the like; cyclo C_{3-6} alkyl encompasses monocyclic alkyl ring structures such as cyclopropyl, cyclobutyl, cyclopentyl, and cyclohexyl; cyclo C_{3-12} alkyl encompasses mono-, bi- or tricyclic alkyl ring structures and is generic to for example cyclopropyl, cyclobutyl, cyclohexyl, cyclohexyl, cyclohectyl, norbornanyl, adamantyl.

The term halo is generic to fluoro, chloro, bromo and iodo. As used in the foregoing and hereinafter, polyhaloC₁₋₆alkyl as a group or part of a group is defined as mono- or polyhalosubstituted C₁₋₆alkyl, in particular methyl with one or more fluoro atoms, for example, difluoromethyl or trifluoromethyl. In case more than one halogen atoms are attached to an alkyl group within the definition of polyhaloC₁₋₆alkyl, they may be the same or different.

When any variable, e.g. aryl, occurs more than one time in any constituent, each definition is independent.

- When any bond is drawn into a ring structure, it means that the corresponding substituent may be linked to any atom of said ring structure. This means for instance that the R¹-C(=X) moiety may be linked to the quinoline or quinolinone moiety in position 5, 6, 7, 8 but also position 3 or position 4.
- For therapeutic use, salts of the compounds of formula (I-A) and (I-B) are those wherein the counterion is pharmaceutically acceptable. However, salts of acids and bases which are non-pharmaceutically acceptable may also find use, for example, in the

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preparation or purification of a pharmaceutically acceptable compound. All salts, whether pharmaceutically acceptable or not are included within the ambit of the present invention.

The pharmaceutically acceptable addition salts as mentioned hereinabove are meant to comprise the therapeutically active non-toxic acid addition salt forms which the compounds of formula (I-A) and (I-B) are able to form. The latter can conveniently be obtained by treating the base form with such appropriate acids as inorganic acids, for example, hydrohalic acids, e.g. hydrochloric, hydrobromic and the like; sulfuric acid; nitric acid; phosphoric acid and the like; or organic acids, for example, acetic, propanoic, hydroxyacetic, 2-hydroxypropanoic, 2-oxopropanoic, oxalic, malonic, succinic, maleic, fumaric, malic, tartaric, 2-hydroxy-1,2,3-propanetricarboxylic, methanesulfonic, ethanesulfonic, benzenesulfonic, 4-methylbenzenesulfonic, cyclohexanesulfamic, 2-hydroxybenzoic, 4-amino-2-hydroxybenzoic and the like acids. Conversely the salt form can be converted by treatment with alkali into the free base form.

The compounds of formula (I-A) and (I-B) containing acidic protons may be converted into their therapeutically active non-toxic metal or amine addition salt forms by treatment with appropriate organic and inorganic bases. Appropriate base salt forms comprise, for example, the ammonium salts, the alkali and earth alkaline metal salts, e.g. the lithium, sodium, potassium, magnesium, calcium salts and the like, salts with organic bases, e.g. primary, secondary and tertiary aliphatic and aromatic amines such as methylamine, ethylamine, propylamine, isopropylamine, the four butylamine isomers, dimethylamine, diethylamine, diethanolamine, dipropylamine, diisopropylamine, di-n-butylamine, pyrrolidine, piperidine, morpholine, trimethylamine, triethylamine, tripropylamine, quinuclidine, pyridine, quinoline and isoquinoline, the benzathine, N-methyl-D-glucamine, 2-amino-2-(hydroxymethyl)-1,3-propanediol, hydrabamine salts, and salts with amino acids such as, for example, arginine, lysine and the like. Conversely the salt form can be converted by treatment with acid into the free acid form.

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The term addition salt also comprises the hydrates and solvent addition forms which the compounds of formula (I-A) and (I-B) are able to form. Examples of such forms are e.g. hydrates, alcoholates and the like.

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The term "quaternary amine" as used hereinbefore defines the quaternary ammonium salts which the compounds of formula (I-A) and (I-B) are able to form by reaction between a basic nitrogen of a compound of formula (I-A) or (I-B) and an appropriate quaternizing agent, such as, for example, an optionally substituted alkylhalide, arylhalide or arylalkylhalide, e.g. methyliodide or benzyliodide. Other reactants with good leaving groups may also be used, such as alkyl trifluoromethanesulfonates, alkyl methanesulfonates, and alkyl p-toluenesulfonates. A quaternary amine has a positively charged nitrogen. Pharmaceutically acceptable counterions include chloro, bromo, iodo, trifluoroacetate and acetate. The counterion of choice can be introduced using ion exchange resins.

It will be appreciated that some of the compounds of formula (I-A) and (I-B) and their *N*-oxides, salts, quaternary amines and stereochemically isomeric forms may contain one or more centers of chirality and exist as stereochemically isomeric forms.

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The term "stereochemically isomeric forms" as used hereinbefore defines all the possible stereoisomeric forms which the compounds of formula (I-A) and (I-B), and their *N*-oxides, salts, quaternary amines or physiologically functional derivatives may possess. Unless otherwise mentioned or indicated, the chemical designation of compounds denotes the mixture of all possible stereoisomeric forms, said mixtures containing all diastereomers and enantiomers of the basic molecular structure as well as each of the individual isomeric forms of formula (I-A) and (I-B) and their *N*-oxides, salts, solvates or quaternary amines substantially free, i.e. associated with less than 10%, preferably less than 5%, in particular less than 2% and most preferably less than 1% of the other isomers. Stereochemically isomeric forms of the compounds of formula (I-A) and (I-B) are obviously intended to be embraced within the scope of the present invention. The same applies to the intermediates as described herein, used to prepare end products of formula (I-A) and (I-B).

The terms *cis* and *trans* are used herein in accordance with Chemical Abstracts nomenclature.

In some compounds of formula (I-A) and (I-B) and in the intermediates used in their preparation, the absolute stereochemical configuration has not been determined. In these cases, the stereoisomeric form which was first isolated is designated as "A" and the second as "B", without further reference to the actual stereochemical configuration. However, said "A" and "B" stereoisomeric forms can be unambiguously characterized

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by physicochemical characteristics such as their optical rotation in case "A" and "B" have an enantiomeric relationship. A person skilled in the art is able to determine the absolute configuration of such compounds using art-known methods such as, for example, X-ray diffraction. In case "A" and "B" are stereoisomeric mixtures, they can be further separated whereby the respective first fractions isolated are designated "A1" and "B1" and the second as "A2" and "B2", without further reference to the actual stereochemical configuration.

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The *N*-oxide forms of the present compounds are meant to comprise the compounds of formula (I-A) and (I-B) wherein one or several nitrogen atoms are oxidized to the so-called *N*-oxide.

Some of the compounds of formula (I-A) and (I-B) may also exist in their tautomeric form. Such forms although not explicitly indicated in the above formula are intended to be included within the scope of the present invention.

Whenever used hereinafter, the term "compounds of formula (I-A) and (I-B)" is meant to also include their *N*-oxide forms, their salts, their quaternary amines and their stereochemically isomeric forms. Of special interest are those compounds of formula (I-A) and (I-B) which are stereochemically pure.

An interesting group of compounds are those compounds of formula (I-A) and (I-B) wherein

- X represents O; $C(R^6)_2$ with R^6 being hydrogen or aryl; or N-R⁷ with R⁷ being amino or hydroxy;
- R¹ represents C₁₋₆alkyl, aryl; thienyl; quinolinyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein the cycloC₃₋₁₂alkyl moiety optionally may contain a double bond and wherein one carbon atom in the cycloC₃₋₁₂alkyl moiety may be replaced by an oxygen atom or an NR⁸-moiety with R⁸ being benzyl or C₁₋₆alkyloxycarbonyl; wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyl, haloC₁₋₆alkyl, hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, halo, aryl, mono— or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, halo, piperazinyl, pyridinyl, morpholinyl, thienyl or a bivalent radical of formula –O- or -O-CH₂-CH₂-O-; or a radical of formula (a-1)

$$Z_1$$
 CH
 Z_2
 $(CH_2)_n$

a-1

wherein Z_1 is a single covalent bond, O or CH_2 ;

Z₂ is a single covalent bond, O or CH₂;

n is an integer of 0, 1, or 2;

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and wherein each hydrogen atom in the phenyl ring independently may optionally be replaced by halo or hydroxy;

or X and R¹ may be taken together with the carbon atom to which X and R¹ are attached to form a radical of formula (b-1), (b-2) or (b-3);

R² represents hydrogen; halo; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkylthio; C₁₋₆alkylcarbonyl; C₁₋₆alkyloxycarbonyl; C₂₋₆alkenyl; hydroxyC₂₋₆alkenyl; C_{2-6} alkynyl; hydroxy C_{2-6} alkynyl; tri $(C_{1-6}$ alkyl)silane C_{2-6} alkynyl; amino; mono— or 15 di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxyC₁₋₆alkyl)amino; mono- or di(C₁₋₆alkylthioC₁₋₆alkyl)amino; aryl; arylC₁₋₆alkyl; arylC₂₋₆alkynyl; C_{1-6} alkyloxy C_{1-6} alkylamino C_{1-6} alkyl; aminocarbonyl optionally substituted with C₁₋₆alkyloxycarbonylC₁₋₆alkyl; a heterocycle selected from thienyl, furanyl, thiazolyl and piperidinyl, optionally 20 N-substituted with morpholinyl or thiomorpholinyl; a radical –NH-C(=0) \mathbb{R}^9 wherein \mathbb{R}^9 represents \mathbb{C}_{1-6} alkyl optionally substituted with cycloC₃₋₁₂alkyl, C₁₋₆alkyloxy, C₁₋₆alkyloxycarbonyl, aryl, aryloxy, thienyl, pyridinyl, mono- or di(C₁₋₆alkyl)amino, C₁₋₆alkylthio, benzylthio, pyridinylthio or pyrimidinylthio; cycloC₃₋₁₂alkyl; cyclohexenyl; amino; arylcycloC₃₋₁₂alkylamino; 25 mono-or-di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxycarbonylC₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxycarbonyl)amino; mono-or di(C₂₋₆alkenyl)amino; mono- or di(arylC₁₋₆alkyl)amino; mono- or diarylamino; arylC₂₋₆alkenyl; furanylC₂₋₆alkenyl; piperididinyl; piperazinyl; indolyl; furyl; benzofuryl; tetrahydrofuryl; indenyl;

adamantyl; pyridinyl; pyrazinyl; aryl or a radical of formula (a-1);

a sulfonamid -NH-SO₂- R^{10} wherein R^{10} represents C_{1-6} alkyl, mono- or poly halo C_{1-6} alkyl, aryl C_{1-6} alkyl or aryl;

- R^3 and R^4 each independently represent hydrogen; $C_{1\text{-}6}$ alkyl; $C_{1\text{-}6}$ alkyloxy $C_{1\text{-}6}$ alkyl; $C_{1\text{-}6}$ alkyloxycarbonyl; or
- R² and R³ may be taken together to form –R²-R³-, which represents a bivalent radical of formula –(CH₂)₄-, –(CH₂)₅-, –Z₄-CH=CH-, –Z₄-CH₂-CH₂-CH₂- or -Z₄-CH₂-CH₂-, with Z₄ being O, S, SO₂ or NR¹¹ wherein R¹¹ is hydrogen, C₁₋₆alkyl, benzyl or C₁₋₆alkyloxycarbonyl; and wherein each bivalent radical is optionally substituted with C₁₋₆alkyl;
- or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH=CH-CH=CH- or -CH₂-CH₂-CH₂-;
 - R⁵ represents hydrogen; piperidinyl; oxo-thienyl; tetrahydrothienyl, arylC₁₋₆alkyl; C₁₋₆alkyloxycarbonylC₁₋₆alkyl or C₁₋₆alkyl optionally substituted with a radical C(=O)NR_xR_y, in which R_x and R_y, each independently are hydrogen, cycloC₃₋₁₂alkyl, C₂₋₆alkynyl or C₁₋₆alkyl optionally substituted with cyano,
- cycloC₃₋₁₂alkyl, C₂₋₆alkynyl or C₁₋₆alkyl optionally substituted with cyan C₁₋₆alkyloxy or C₁₋₆alkyloxycarbonyl;
 - Y represents O or S;
 - or Y and R⁵ may be taken together to form =Y-R⁵- which represents a radical of formula

20 -CH=N-N= (c-1); or

-N=N-N= (c-2);

- aryl represents phenyl or naphthyl optionally substituted with one or more substituents selected from halo, C₁₋₆alkyloxy, phenyloxy, mono-or di(C₁₋₆alkyl)amino and cyano;
- and when the R¹-C(=X) moiety is linked to another position than the 7 or 8 position, then said 7 and 8 position may be substituted with R¹⁵ and R¹⁶ wherein either one or both of R¹⁵ and R¹⁶ represents C₁₋₆alkyl or R¹⁵ and R¹⁶ taken together may form a bivalent radical of formula -CH=CH-CH=CH-.
- A further most interesting group of compounds comprises those compounds of formula (I-A) and (I-B) wherein X represents O;
 - R¹ represents C₁₋₆alkyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyloxy, aryl, halo or thienyl;
- R² represents hydrogen; halo; C₁₋₆alkyl or amino; R³ and R⁴ each independently represent hydrogen or C₁₋₆alkyl; or

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 R^2 and R^3 may be taken together to form $-R^2-R^3$ -, which represents a bivalent radical of formula $-Z_4$ - CH_2 - CH_2 - CH_2 - or $-Z_4$ - CH_2 - CH_2 - with Z_4 being O or NR^{11} wherein R^{11} is C_{1-6} alkyl; and wherein each bivalent radical is optionally substituted with C_{1-6} alkyl;

or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH₂-CH₂-CH₂-CH₂-;

R⁵ represents hydrogen;

Y represents O; and

aryl represents phenyl optionally substituted with halo.

A further interesting group of compounds comprises those compounds of formula (I-A) and (I-B) wherein the R¹-C(=X) moiety is linked to the quinoline or quinolinone moiety in position 6.

In order to simplify the structural representation of some of the present compounds and intermediates in the following preparation procedures, the quinoline or the quinolinene moiety will hereinafter be represented by the symbol Q.

$$Q = \bigvee_{N \to \mathbb{R}^2}^{\mathbb{R}^4} \qquad \text{or} \qquad \bigvee_{N \to \mathbb{R}^5}^{\mathbb{R}^4}$$

The compounds of formula (I-A) or (I-B), wherein X represents O, said compounds being represented by formula ($I_{A/B}$ -a), can be prepared by oxidizing an intermediate of formula (II) in the presence of a suitable oxidizing agent, such as potassium permanganate, and a suitable phase-transfer catalyst, such as tris(dioxa-3,6-heptyl)amine, in a suitable reaction-inert solvent, such as for example dichloromethane.

$$R^{1}$$
— CH — Q oxidation R^{1} — C — Q

(II) $(I_{A/B}$ - $a)$

Compounds of formula (I_{A/B}-a) may also be prepared by reacting an intermediate of formula (III) with an intermediate of formula (IV), wherein W₁ represents a halo atom, e.g. bromo, in the presence of butyl lithium and a suitable reaction-inert solvent, such as for example tetrahydrofuran.

$$R^{1} - C = N + W_{1} - Q \longrightarrow R^{1} - C - Q$$

$$(III) \qquad (IV) \qquad (I_{A/B} - a)$$

Alternatively, compounds of formula ($I_{A/B}$ -a) may also be prepared by reacting an intermediate of formula (IV) with an intermediate of formula (IV) in the presence of butyl lithium and a suitable reaction-inert solvent, such as for example tetrahydrofuran.

$$R^{1} = C - N CH_{3}$$
 + $W_{1} = Q$ \longrightarrow $R^{1} = C - Q$ (IV) (I_{A/B}-a)

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Compounds of formula ($I_{A/B}$ -a), wherein the R^1 substituent is linked to the carbonyl moiety via an oxygen atom, said R^1 substituent being represented by O- R^{1a} and said compounds by formula ($I_{A/B}$ -a-1), can be prepared by reacting an intermediate of formula (VI) with an intermediate of formula (VII) in the presence of a suitable acid, such as sulfuric acid.

$$R^{1a}$$
 OH + HO C Q \rightarrow R^{1a} O C Q (VI) ($I_{A/B}$ -a-1)

Compounds of formula (I-A), wherein R² represents methylcarbonyl, said compounds being represented by formula (I-A-1), can be prepared by reacting an intermediate of formula (VIII) in the presence of a suitable acid, such as hydrochloric acid, and a suitable reaction-inert solvent, such as for example tetrahydrofuran.

The compounds of formula (I) may also be converted into each other following art-known transformations.

Compounds of formula (I-A) wherein R² is a halo atom, such as chloro, can be converted into a compound of formula (I-A), wherein R² is another halo atom, such as fluoro or iodo, by reaction with a suitable halogenating agent, such as for example

potassium fluoride or sodium iodide, in the presence of a suitable reaction-inert solvent, e.g. dimethyl sulfoxide or acetonitrile and optionally in the presence of acetyl chloride.

Compounds of formula (I-A), wherein R^2 is a suitable leaving group, such as a halo atom, e.g. chloro, iodo, said leaving group being represented by W^2 and said compounds by (I-A-2), can be converted into a compound of formula (I-A) wherein R^2 is cyano, said compound being represented by formula (I-A-3), by reaction with a suitable cyano-introducing agent, such as for example trimethylsilanecarbonitrile, in the presence of a suitable base such as N,N-diethylethanamine and a suitable catalyst, such as for example tetrakis(triphenylphosphine)palladium.

Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A-4) by reaction with C_{2-6} alkynyltri(C_{1-6} alkyl)silane in the presence of CuI, an appropriate base, such as for example N,N-diethylethanamine, and an appropriate catalyst, such as for example tetrakis(triphenylphosphine)palladium. Compounds of formula (I-A-4) can on their turn be converted into a compound of formula (I-A-5) by reaction with potassium fluoride in the presence of a suitable acid such as acetic acid, or by reaction with a suitable base, such as potassium hydroxide, in the presence of a suitable reaction-inert solvent, such as an alcohol, e.g. methanol and the like.

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Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A-6) by reaction with an intermediate of formula (IX) in the presence of CuI, a suitable base, such as for example *N*,*N*-diethylethanamine, and a suitable catalyst such as tetrakis(triphenylphosphine)palladium.

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Compounds of formula (I-A-2) can also be converted into a compound wherein R^2 is C_{1-6} alkyl, said compound being represented by formula (I-A-8) in the presence of a suitable alkylating agent, such as for example $Sn(C_{1-6}$ alkyl)₄, or into a compound wherein R^2 is C_{2-6} alkenyl, said compound being represented by formula (I-A-9) in the presence of a suitable alkenylating agent, such as for example $Sn(C_{2-6}$ alkenyl)(C_{1-6} alkyl)₃, both reactions in the presence of a suitable catalyst, such as for example tetrakis(triphenylphosphine)palladium and a reaction-inert solvent, such as for example toluene or dioxane.

Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A-7) wherein Z represents O or S, by reaction with an intermediate of formula (X)

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optionally in the presence of a suitable base such as dipotassium carbonate and a reaction-inert solvent, such as *N*,*N*-dimethyl formamide.

Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A), wherein R² is C₁₋₆alkyloxycarbonyl, said compound being represented by formula (I-A-10) and a compound of formula (I-A), wherein R² is hydrogen, said compound being represented by formula (I-A-11), by reaction with a suitable alcohol of formula

C₁₋₆alkylOH and CO in the presence of a suitable catalyst, such as for example palladium(II)acetate, triphenylphosphine, a suitable base such as dipotassium carbonate and a reaction-inert solvent, such as *N*,*N*-dimethylformamide.

Compounds of formula (I-A-11) can also be prepared by reacting a compound of formula (I-A-2) with Zn in the presence of a suitable acid such as acetic acid.

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Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A), wherein R^2 is aminocarbonyl substituted with C_{1-6} alkyloxycarbonyl C_{1-6} alkyl, said compound being represented by formula (I-A-12), by reaction with an intermediate of formula H_2N-C_{1-6} alkyl- $C(=O)-O-C_{1-6}$ alkyl in the presence of CO, a suitable catalyst such as tetrakis(triphenylphosphine)palladium, a suitable base, such as for example N_1N -diethylethanamine, and a suitable reaction-inert solvent, such as for example toluene.

$$\begin{array}{c} \mathbb{R}^1 \\ \mathbb{R}^4 \\ \mathbb{R}^3 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^5 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^5 \\ \mathbb{R}^4 \\ \mathbb{R}^5 \\ \mathbb{R}^$$

Compounds of formula (I-A-2) can also be converted into a compound of formula (I-A) wherein R² is aryl or a heterocycle selected from the group described in the definition of R² hereinabove, said R² being represented by R^{2a} and said compound by formula (I-A-13) by reaction with an intermediate of formula (XI), (XII) or (XIII) in the presence of a suitable catalyst such as for example tetrakis(triphenylphosphine)palladium and a suitable reaction-inert solvent, such as for example dioxane.

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Compounds of formula (I-A-2) can also be converted into a compound of formula (I-B), wherein Y and R⁵ are taken together to form a radical of formula (b-1) or (b-2), said compound being represented by formula (I-B-1) or (I-B-2), by reaction with hydrazincarboxaldehyde or sodium azide in a suitable reaction-inert solvent, such as an alcohol, e.g. butanol, or *N*,*N*-dimethylformamide.

$$R^{1} \longrightarrow R^{2}$$

$$R^{2} \longrightarrow R^{3}$$

$$(I-B-1)$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{1} \longrightarrow R^{4}$$

$$R^{2} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{4} \longrightarrow R^{4}$$

$$R^{4} \longrightarrow R^{4}$$

$$R^{4} \longrightarrow R^{4}$$

$$R^{3} \longrightarrow R^{4}$$

$$R^{4} \longrightarrow R^{4}$$

$$R^{5} \longrightarrow R^{4}$$

$$R^$$

Compounds of formula (I-A-11) can be converted into the corresponding N-oxide, represented by formula (I-A-14), by reaction with a suitable peroxide, such as 3-chloro-benzenecarboperoxoic acid, in a suitable reaction-inert solvent, such as for

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example methylene chloride. Said compound of formula (I-A-14) can further be converted into a compound of formula (I-B), wherein R⁵ is hydrogen, said compound being represented by formula (I-B-3), by reaction with 4-methyl-benzene sulfonyl chloride in the presence of a suitable base, such as for example dipotassium carbonate and a suitable reaction-inert solvent, such as for example methylene chloride.

(I-A-11)
$$\stackrel{\text{peroxide}}{\longrightarrow}$$
 $\stackrel{\text{R}^4}{\longrightarrow}$ $\stackrel{\text{R$

Compounds of formula (I-B-3) can also be prepared from a compound of formula (I-A), wherein R^2 is C_{1-6} alkyloxy, said compound being represented by formula (I-A-15), by reaction with a suitable acid, such as hydrochloric acid, in the presence of a suitable reaction-inert solvent, such as for example tetrahydrofuran.

$$R^{1}$$
 R^{2}
 R^{2}
 R^{3}
 R^{1}
 R^{2}
 R^{3}
 R^{1}
 R^{2}
 R^{3}
 R^{1}
 R^{2}
 R^{3}
 R^{4}
 R^{3}
 R^{1}
 R^{2}
 R^{3}
 R^{4}
 R^{4}
 R^{3}
 R^{4}
 R^{4}
 R^{3}
 R^{4}
 R^{4

15 Compounds of formula (I-B-3) can be converted into a compound of formula (I-B), wherein R⁵ represents C₁₋₆alkyl, said compound being represented by formula (I-B-4), by reaction with an appropriate alkylating agent, such as for example an intermediate of formula (XIV), wherein W₃ represents a suitable leaving group such as a halo atom e.g. iodo, in the presence of potassium tert. butoxide and in the presence of a suitable reaction-inert solvent, such as for example tetrahydrofuran.

$$R^{1}$$
 R^{1}
 R^{2}
 R^{3}
 R^{1}
 R^{3}
 R^{4}
 R^{5}
 R^{4}
 R^{5}
 R^{4}
 R^{5}
 R^{5}
 R^{6}
 R^{6}
 R^{7}
 R^{7

Compounds of formula (I-B-3) can also be converted into a compound of formula (I-B), wherein R^5 is C_{1-6} alkyloxycarbonyl C_{1-6} alkyl or aryl C_{1-6} alkyl, said R^5 being represented by R^{5a} and said compound being represented by formula (I-B-5), by reaction with an intermediate of formula (XV), wherein W_4 represents a suitable leaving group, such as a halo atom, e.g. bromo, chloro and the like, in the presence of a suitable base, such as for example sodium hydride and a suitable reaction-inert solvent, such as for example N,N-dimethylformamide.

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$$R^{1}$$
 R^{1}
 R^{2}
 R^{3}
 R^{5a}
 R^{4}
 R^{3}
 R^{5a}
 R^{5a}

Compounds of formula (I-A-2) can also be converted into a compound of formula (I-B), wherein R⁵ is hydrogen and Y is S, said compound being represented by formula (I-B-6), by reaction with H₂N-C(=S)-NH₂ in the presence of a suitable base, such as potassium hydroxide, and a suitable reaction-inert solvent, such as an alcohol, for example ethanol, or water. Compounds of formula (I-B-6) can further be converted

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into a compound of formula (I-A), wherein R^2 is C_{1-6} alkylthio, said compound being represented by formula (I-A-16), by reaction with a suitable C_{1-6} alkylhalide, such as for example C_{1-6} alkyliodide, in the presence of a suitable base, such as dipotassium carbonate, and a suitable solvent, such as for example acetone.

$$R^{1} - C - NH_{2} + H_{2}N - C - NH_{2}N - C - NH_{2} + H_{2}N - C - NH_{2}N - C - NH_{2}N - C - NH_{2}N - C - NH_{2}N - C - NH_{$$

Compounds of formula ($I_{A/B}$ -a) can be converted into a compounds of formula ($I_{A/B}$ -b) or (I_{-B}), wherein X is N-R⁷, said compound being represented by formula ($I_{A/B}$ -b), by reaction with an intermediate of formula (XVI), optionally in the presence of a suitable base, such as for example $N_{,N}$ -diethylethanamine, and in the presence of a suitable reaction-inert solvent, such as an alcohol, e.g. ethanol.

$$R^{1} - C - Q + R^{7} - NH_{2} - R^{1} - C - Q$$

$$(I_{A/B}-a) \qquad (XVI) \qquad (I_{A/B}-b)$$

As already indicated in the preparation procedure of compounds of formula (I-A-13) described above, the compounds of formula (I) may also be converted to the corresponding *N*-oxide forms following art-known procedures for converting a trivalent nitrogen into its *N*-oxide form. Said *N*-oxidation reaction may generally be carried out by reacting the starting material of formula (I) with an appropriate organic or inorganic peroxide. Appropriate inorganic peroxides comprise, for example, hydrogen peroxide, alkali metal or earth alkaline metal peroxides, e.g. sodium peroxide, potassium peroxide; appropriate organic peroxides may comprise peroxy acids such as, for example, benzenecarboperoxoic acid or halo substituted benzenecarboperoxoic acid, e.g. 3-chlorobenzenecarboperoxoic acid, peroxoalkanoic acids, e.g. peroxoacetic acid, alkylhydroperoxides, e.g. tert-butyl hydroperoxide. Suitable solvents are, for example, water, lower alkanols, e.g. ethanol and the like, hydrocarbons, e.g. toluene, ketones, e.g. 2-butanone, halogenated hydrocarbons, e.g. dichloromethane, and mixtures of such solvents.

Some of the intermediates and starting materials used in the above reaction procedures are commercially available, or may be synthesized according to procedures already described in the literature.

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Intermediates of formula (II) may be prepared by reacting an intermediate of formula (XVII) with an intermediate of formula (XVIII), wherein W_5 represents a suitable leaving group such as a halo atom, e.g. chloro, bromo and the like, in the presence of magnesium, diethylether and a suitable reaction-inert solvent, such as diethylether.

$$Q = C + H + H^{1} = W_{5} \longrightarrow H^{1} = CH = Q$$
(XVIII) (III)

Intermediates of formula (XVII) may be prepared by oxidizing an intermediate of formula (XIX) in the presence of a suitable oxidizing agent, such as MnO₂, and a suitable reaction-inert solvent, such as methylene chloride.

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Intermediates of formula (XIX) can be prepared by reducing an intermediate of formula (XX) in the presence of a suitable reducing agent such as lithium aluminium hydride, and a suitable reaction-inert solvent, such as tetrahydrofuran.

$$\begin{array}{c} O \\ \square \\ Q - C - O - C_{1^-6} \text{alkyl} \end{array} \qquad \begin{array}{c} \text{reduction} \\ \longrightarrow Q - C \text{H}_2 - O \text{H} \end{array}$$
(XIX)

Intermediates of formula (XX), wherein Q represents a quinoline moiety optionally substituted in position 3 with C₁₋₆alkyl and wherein the carbonyl moiety is placed in position 6, said intermediates being represented by formula (XX-a), can be prepared by reacting an intermediate of formula (XXI) with an intermediate of formula (XXII) in the presence of sodium 3-nitro-benzene sulfonate, a suitable acid, such as sulfuric acid, and a suitable alcohol, e.g. methanol, ethanol, propanol, butanol and the like.

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

Alternatively, intermediates of formula (II) can also be prepared by reacting an intermediate of formula (XXIII) with an intermediate of formula (XXIV), wherein W_6 is a suitable leaving group, such as a halo atom, e.g. bromo, chloro and the like, in the presence of a suitable agent, such as butyl lithium and a suitable reaction-inert solvent, such as tetrahydrofuran.

$$R^{1} \xrightarrow{C} H + Q \xrightarrow{W_{6}} R^{1} \xrightarrow{CH-Q}$$
(XXIII) (XXIV) (II)

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Intermediates of formula (XXIII) can be prepared by oxidizing an intermediate of formula (XXV) using the Moffatt Pfitzner or Swern oxidation (dimethylsulfoxide adducts with dehydrating agents e.g. DCC, Ac₂O, SO₃, P₄O₁₀, COCl₂ or Cl-CO-COCl) in an inert solvent such as methylene chloride.

$$R^{1}$$
 CH_{2} OH R^{1} C H (XXV) $(XXIII)$

Intermediates of formula (XXV) can be prepared by reducing an intermediate of formula (XXVI) in the presence of a suitable reducing agent, such as for example lithium aluminium hydride and a suitable reaction-inert solvent, such as benzene.

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$$R^{1} = C = O - C_{1-6} \text{alkyl} \qquad \Rightarrow \qquad R^{1} = CH_{2} - OH$$
(XXVI) (XXV)

Intermediates of formula (XXVI) can be prepared from an intermediate of formula (XXVII) by esterification in the presence of a suitable alcohol, such as methanol, ethanol, propanol, butanol and he like, and a suitable acid, such as sulfuric acid.

Intermediates of formula (XXVII), wherein R^1 represents a radical of formula (a-1) with Z_1 being O, Z_2 being CH_2 and n being 1, said intermediates being represented by formula (XXVII-a), can be prepared by reducing an intermediate of formula (XXVIII) in the presence of a suitable reducing agent such as hydrogen, and a suitable catalyst, such as palladium on charcoal, and a suitable acid such as acetic acid. When R^1 of intermediate (XXVII) represents an optionally substituted phenyl moiety, it can also be converted into an optionally substituted cyclohexyl moiety by reduction in the presence of a suitable reducing agent such as rhodium on Al_2O_3 , and a suitable reaction-inert solvent, such as tetrahydrofuran.

Intermediates of formula (IV), wherein Q represents a quinoline moiety substituted in position 2 with halo,e.g. chloro, said intermediates being represented by formula (IV-a), can be prepared by reacting an intermediate of formula (IV), wherein Q represents a quinolinone moiety with R⁵ being hydrogen, said intermediate being represented by formula (IV-b), in the presence of POCl₃.

$$\begin{array}{c}
W_1 \\
\downarrow \\
N \\
\downarrow \\
N
\end{array}$$
(IV-a)

Intermediates of formula (IV-a), wherein R⁴ is hydrogen, said intermediates being represented by formula (IV-a-1), can also be prepared by reacting an intermediate of formula (XXIX) with POCl₃ in the presence of *N*,*N*-dimethylformamide (Vilsmeier-Haack formylation followed by cyclization).

$$\begin{array}{c} W_1 \\ O \\ NH - C \\ R^3 \end{array}$$

$$(IV-a-1)$$

Intermediates of formula (XXIX) may be prepared by reacting an intermediate of formula (XXX) with an intermediate of formula (XXXI), wherein W_7 represents a suitable leaving group, such as a halo atom, e.g. chloro, in the presence of a suitable base, such as for example N,N-diethylethanamine, and a suitable reaction-inert solvent, such as methylene chloride.

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Intermediates of formula (IV-a) can be converted into an intermediate of formula (IV-c) by reaction with an intermediate of formula (XXXII) in the presence of a suitable reaction-inert solvent, such as an alcohol, e.g. methanol and the like.

Intermediates of formula (IV-a) can also be converted into an intermediate of formula (IV-d-1) by reaction with a suitable amine of formula (XXXIII-a), wherein Z_3 and Z_4 each independently represent hydrogen, $C_{1\text{-}6}$ alkyl, $C_{1\text{-}6}$ alkyloxy $C_{1\text{-}6}$ alkyl,

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 C_{1-6} alkylthio C_{1-6} alkyl or into an intermediate of formula (IV-d-2) by reaction with a suitable amine of formula (XXXIII-b), wherein Z_3 and Z_4 are taken together to form a heterocycle as defined hereinabove in the definition of R^2 provided that the heterocycle comprises at least one nitrogen atom, in the presence of a suitable base, such as for example dipotassium carbonate, and a reaction-inert solvent, such as N,N-dimethylformamide.

Intermediates of formula (IV-a), wherein R³ represents CH₂-CH₂-CH₂-Cl, said intermediates being represented by formula (IV-a-2), can also be converted into an intermediate of formula (IV), wherein R² and R³ are taken together to form a bivalent radical of formula –O-CH₂-CH₂-CH₂-, said intermediate being represented by formula (IV-e-1), by reaction with a suitable acid, such as hydrochloric acid and the like. Intermediates of formula (IV-a-2) can also be converted into an intermediate of formula (IV), wherein R² and R³ are taken together to form a bivalent radical of formula

-S-CH₂-CH₂-, said intermediate being represented by formula (IV-e-2), by reaction with H₂N-C(=S)-NH₂ in the presence of a suitable reaction-inert solvent, such as an alcohol, e.g. ethanol.

Intermediates of formula (V) may be prepared by reacting an intermediate of formula (XXVII) with an intermediate of formula CH₃-NH-O-CH₃ in the presence of 1,1'-carbonyldiimidazole and a suitable reaction-inert solvent, such as methylene chloride.

$$R^{1}$$
 C OH $+$ $H_{3}C$ $-NH$ $-O$ $-CH_{3}$ $-CH_{3}$ $-CH_{3}$ $-CH_{3}$ $-CH_{3}$ $-CH_{3}$

Intermediates of formula (VII), wherein Q represents a quinoline moiety, in particular a quinoline moiety wherein R² is ethyl, R³ is methyl and R⁴ is hydrogen, and the carboxyl moiety is placed in position 6, said intermediates being represented by formula (VII-a), can be prepared by reaction an intermediate of formula (XXXIV) in the presence of a suitable aldehyde, such as CH₃-CH₂-CH(=O), (CH₂O)_n, ZnCl₂, FeCl₃ and a suitable reaction-inert solvent, such as an alcohol, for example ethanol.

$$CH_3$$
- CH_2 - $CH(=O)$
 $(CH_2O)_n$
 $(VII-a)$

Intermediates of formula (VIII) can be prepared by reacting an intermediate of formula (XXXV) with an intermediate of formula (XXXVI) in the presence of a suitable

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catalyst, such as for example tetrakis(triphenylphosphine)palladium and a suitable reaction-inert solvent, such as for example dioxane.

Still some other preparations can be devised, some of them are disclosed further in this application with the Examples.

- Pure stereoisomeric forms of the compounds and the intermediates of this invention 10 may be obtained by the application of art-known procedures. Diastereomers may be separated by physical separation methods such as selective crystallization and chromatographic techniques, e.g. liquid chromatography using chiral stationary phases. Enantiomers may be separated from each other by the selective crystallization of their diastereomeric salts with optically active acids. Alternatively, enantiomers may be 15 separated by chromato-graphic techniques using chiral stationary phases. Said pure stereoisomeric forms may also be derived from the corresponding pure stereoisomeric forms of the appropriate starting materials, provided that the reaction occurs stereoselectively or stereospecifically. Preferably, if a specific stereoisomer is desired, said compound will be synthesized by stereoselective or stereospecific methods of 20 preparation. These methods will advantageously employ chirally pure starting materials. Stereoisomeric forms of the compounds of formula (I) are obviously intended to be included within the scope of the invention.
- A stereoisomer of a compound of formula (I-A) or (I-B) such as a *cis* form, may be converted into another stereoisomer such as the corresponding *trans* form by reacting the compound with a suitable acid, such as hydrochloric acid, in the presence of a suitable reaction-inert solvent, such as for example tetrahydrofuran.
- The compounds of formula (I-A) and (I-B), the *N*-oxides, the pharmaceutically acceptable addition salts, the quaternary amines and the stereochemically isomeric forms thereof show mGluR antagonistic activity, more in particular Group I mGluR

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antagonistic activity. The Group I mGluR specifically antagonized by the present compounds is the mGluR1.

The mGluR1 antagonistic activity of the present compounds can be demonstrated in the Signal transduction on cloned rat mGluR1 in CHO cells test and the Cold allodynia test in rats with a Bennett ligation, as described hereinafter.

Due to their mGluR antagonistic activity, more in particular their Group I mGluR antagonistic activity and even more in particular, their mGluR1 antagonistic activity, the compounds of formula (I-A) or (I-B), their N-oxides, pharmaceutically acceptable addition salts, quaternary amines and stereochemically isomeric forms are useful in the treatment or prevention of glutamate-induced diseases of the central nervous sytem. Diseases in which a role for glutamate has been demonstrated include drug addiction or abstinence (dependence, opioid tolerance, opioid withdrawal), hypoxic, anoxic and ischemic injuries (ischemic stroke, cardiac arrest), pain (neuropathic pain, inflammatory pain, hyperalgesia), hypoglycemia, diseases related to neuronal damage, brain trauma, head trauma, spinal cord injury, myelopathy, dementia, anxiety, schizophrenia, depression, impaired cognition, amnesia, bipolar disorders, conduct disorders, Alzheimer's disease, vascular dementia, mixed (Alzheimer's and vascular) dementia, Lewy Body disease, delirium or confusion, Parkinson's disease, Huntington's disease, Down syndrome, epilepsy, aging, Amyotrophic Lateral Sclerosis, multiple sclerosis, AIDS (Acquired Immune Deficiency Syndrome) and AIDS related complex (ARC).

In view of the utility of the compounds of formula (I-A) and (I-B), there is provided a method of treating warm-blooded animals, including humans, suffering from glutamate-induced diseases of the central nervous system. Said method comprises the administration, preferably oral administration, of an effective amount of a compound of formula (I-A) or (I-B), a *N*-oxide form, a pharmaceutically acceptable addition salt, a quaternary amine or a possible stereoisomeric form thereof, to warm-blooded animals, including humans.

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In view of the above described pharmacological properties, the compounds of formula (I-A) and (I-B) or any subgroup thereof, their *N*-oxides, pharmaceutically acceptable addition salts, quaternary amines and stereochemically isomeric forms, may be used as a medicine. In particular, the use of a compound of formula (I-A) and (I-B) in the manufacture of a medicament for treating or preventing glutamate-induced diseases of

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the central nervous system is provided. More in particular, the present compounds may be used as neuroprotectants, analgesics or anticonvulstants.

The present invention also provides compositions for treating or preventing glutamate-induced diseases of the central nervous system comprising a therapeutically effective amount of a compound of formula (I-A) or (I-B) and a pharmaceutically acceptable carrier or diluent.

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Therefore, the compounds of the present invention or any subgroup thereof may be formulated into various pharmaceutical forms for administration purposes. As appropriate compositions there may be cited all compositions usually employed for systemically administering drugs. To prepare the pharmaceutical compositions of this invention, a therapeutically effective amount of a particular compound, in base or addition salt form, as the active ingredient is combined in intimate admixture with a pharmaceutically acceptable carrier, which carrier may take a wide variety of forms depending on the form of preparation desired for administration. These pharmaceutical compositions are desirably in unitary dosage form suitable, preferably, for administration orally, rectally, topically, percutaneously or by parenteral injection. For example, in preparing the compositions in oral dosage form, any of the usual pharmaceutical media may be employed, such as, for example, water, glycols, oils, alcohols and the like in the case of oral liquid preparations such as suspensions, syrups, emulsions, elixirs and solutions: or solid carriers such as starches, sugars, kaolin, lubricants, binders, disintegrating agents and the like in the case of powders, pills, capsules and tablets. Because of their ease in administration, tablets and capsules represent the most advantageous oral dosage unit form, in which case solid pharmaceutical carriers are obviously employed. For parenteral compositions, the carrier will usually comprise sterile water, at least in large part, though other ingredients, for example, to aid solubility, may be included. Injectable solutions, for example, may be prepared in which the carrier comprises saline solution, glucose solution or a mixture of saline and glucose solution. Injectable suspensions may also be prepared in which case appropriate liquid carriers, suspending agents and the like may be employed. Also included are solid form preparations which are intended to be converted, shortly before use, to liquid form preparations. As appropriate compositions for topical application there may be cited all compositions usually employed for topically administering drugs e.g. creams, gel, dressings, shampoos, tinctures, pastes, ointments, salves, powders and the like. In the compositions suitable for percutaneous administration, the carrier optionally comprises a penetration enhancing agent and/or a

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suitable wetting agent, optionally combined with suitable additives of any nature in minor proportions, which additives do not cause a significant deleterious effect to the skin. Said additives may facilitate the administration to the skin and/or may be helpful for preparing the desired compositions. These compositions may be administered in various ways, e.g., as a transdermal patch, as a spot-on, as an ointment.

It is especially advantageous to formulate the aforementioned pharmaceutical compositions in unit dosage form for ease of administration and uniformity of dosage. Unit dosage form as used in the specification and claims herein refers to physically discrete units suitable as unitary dosages, each unit containing a predetermined quantity of active ingredient calculated to produce the desired therapeutic effect in association with the required pharmaceutical carrier. Examples of such unit dosage forms are tablets (including scored or coated tablets), capsules, pills, suppositories, powder packets, wafers, injectable solutions or suspensions, teaspoonfuls, tablespoonfuls and the like, and segregated multiples thereof.

The therapeutically effective dose or frequency of administration depends on the particular compound of formula (I-A) or (I-B) used, the particular condition being treated, the severity of the condition being treated, the age, weight, sex, fed or fasted state, and the general physical condition of the particular patient as well as other medication the individual may be taking, as is well known to those skilled in the art. Furthermore, it is evident that said therapeutically effective dose or the effective daily dose may be lowered or increased depending on the response of the treated subject and/or depending on the evaluation of the physician prescribing the compounds of the instant invention. It may be appropriate to administer the required dose as two, three, four or more sub-doses at appropriate intervals throughout the day. Said sub-doses may be formulated as unit dosage forms.

The following examples are intended to illustrate the present invention.

Experimental part

Hereinafter, "DMF" is defined as *N*,*N*-dimethylformamide, "DIPE" is defined as diisopropylether, "DMSO" is defined as dimethylsulfoxide, "BHT" is defined as 2,6-bis(1,1-dimethylethyl)-4-methylphenol, and "THF" is defined as tetrahydrofuran.

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Preparation of the intermediates

Example A1

Preparation of

A mixture of 4-(1-methylethoxy)benzoic acid (0.083 mol) and Rh/Al₂O₃ 5% (10g) in THF (220ml) was hydrogenated at 50°C (under 3 bar pressure of H₂) for 1 night. The mixture was filtered over celite, washed with THF and evaporated. Yield: 16g of intermediate 1 (100%).

Example A2

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Preparation of 2-ethyl-3-methyl-6-quinolinecarboxylic acid (interm. 2)

A mixture of 4-aminobenzoic acid (0.299 mol) in ethanol (250ml) was stirred at room temperature. ZnCl₂ (0.0367 mol) and (CH₂O)_n (10g) were added. FeCl₃.6H₂O (0.5 mol) was added portionwise and the temperature rised till 60-65°C. Propanal (30ml) was added dropwise over a 2 hours period. The mixture was refluxed for 2 hours and kept at room temperature for 12 hours. The mixture was poured into water and filtered through celite. The filtrate was acidified till pH=7 with HCl 6N and the mixture was evaporated till dryness. The residue was used without further purification. Yield: 56.1g of 2-ethyl-3-methyl-6-quinolinecarboxylic acid (interm. 2).

Example A3

Preparation of

Pentanoyl chloride (0.2784 mol) was added at 5°C to a mixture of 4-bromobenzenamine (0.232 mol) and *N*,*N*-diethylethanamine (0.2784 mol) in CH₂Cl₂ (400ml). The mixture was stirred at room temperature overnight, poured out into water and extracted with CH₂Cl₂. The organic layer was separated, washed with a concentrated NH₄OH solution and water, dried (MgSO₄), filtered and the solvent was evaporated. The residue (60g) was crystallized from diethylether. The precipitate was filtered off and dried. The residue (35g, 63%) was taken up in CH₂Cl₂. The organic layer was separated, washed with a 10% K₂CO₃ solution, washed with water, dried (MgSO₄), filtered and the solvent was evaporated. Yield: 30g of intermediate (3) (54%).

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Example A4

Preparation of NCI (interm. 4)

A mixture of 6-bromo-2(1*H*)-quinolinone (0.089 mol) in POCl₃ (55ml) was stirred at 60°C overnight, then at 100°C for 3 hours and the solvent was evaporated. The residue was taken up in CH₂Cl₂, poured out into ice water, basified with NH₄OH conc., filtered over celite and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. Yield: 14.5g of intermediate (4) (67%).

Example A5

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a) Preparation of CINN (interm. 5)

DMF (37ml) was added dropwise at 10°C under N₂ flow to POCl₃ (108ml). After complete addition, the mixture was allowed to warm to room temperature. N-(4-bromophenyl)butanamide (0.33 mol) was added portionwise. The mixture was stirred at 85°C overnight, then allowed to cool to room temperature and poured out on ice (exothermic reaction). The precipitate was filtered off, washed with a small amount of water and dried (vacuum). The residue was washed with EtOAc/diethyl ether and dried. Yield: 44.2g of intermediate (5) (50%).

A mixture of intermediate (5) (0.162 mol) in methanol (600ml), and a solution of methanol sodium salt in methanol at 35% (154ml) was stirred and refluxed overnight. The mixture was poured out on ice. The precipitate was filtered off, washed with a small amount of water and taken up in CH₂Cl₂. K₂CO₃ 10% was added and the mixture was extracted with CH₂Cl₂. The organic layer was separated, washed with water, dried (MgSO₄), filtered and the solvent was evaporated. Yield: 31.9g of intermediate (6) (74%).

Example A6

Preparation of (interm. 7)

1,1'-Carbonylbis-1*H*-imidazole (0.074 mol) was added portionwise to a mixture of 4-methoxycyclohexanecarboxylic acid (0.063 mol) in CH₂Cl₂ (200ml). The mixture was stirred at room temperature for 1 hour. Then *N*-methoxymethanamine (0.074 mol)

was added. The mixture was stirred at room temperature overnight, poured out into H_2O and extracted with CH_2Cl_2 . The organic layer was separated, washed several times with H_2O , dried (MgSO₄), filtered and the solvent was evaporated. Yield: 12.6g of interm. 7.

5 Example A7

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10) (85%).

- a) A mixture of 6-fluoro-4-oxo-4H-1-benzopyran-2-carboxylic acid (0.30mol) in acetic acid (400ml) was hydrogenated with Pd/C (3g) as a catalyst. After uptake of H_2 (3 equiv), the catalyst was filtered off. The filtrate was evaporated. The residue was stirred in petroleum ether. The precipitate was filtered off and dried (vacuum; 70°C).
- After recrystallization from CHCl₃/CH₃OH, the precipitate was filtered off and dried (vacuum; 80°C and high vacuum; 85°C). Yield: 8.8 g of 6-fluoro-3,4-dihydro-2*H*-1-benzopyran-2-carboxylic acid (interm. 8) (15.0%).
 - b) A mixture of intermediate (8) (0.255 mol) in ethanol (400ml) and H₂SO₄ (5ml) was stirred and refluxed for 8 hours. The solvent was evaporated till dryness. The residue was dissolved in CH₂Cl₂. The organic layer was separated, washed with K₂CO₃ 10%, dried (MgSO₄), filtered and the solvent was evaporated. Yield: 45g of ethyl 6-fluoro-3,4-dihydro-2*H*-1-benzopyran-2-carboxylate (interm. 9) (79%).
 - c) Reaction under N₂. A mixture of sodium bis(2-methoxyethoxy)aluminumhydride, 70 wt % solution in methylbenzene 3.4M (0.44 mol) in benzene (150 ml) (reflux) was added dropwise during 1 hour to a refluxed mixture of interm. 9 (0.22 mol) and benzene (600 ml). After stirring for 2.5 hours at reflux temperature, the mixture was cooled to ±15°C. The mixture was decomposed by adding dropwise ethanol (30 ml) and water (10 ml). This mixture was poured out onto ice/water and this mixture was acidified with concentrated hydrochloric acid. This mixture was extracted with diethyl ether (500 ml). The separated organic layer was washed with water, dried, filtered and the solvent was evaporated. The residue was purified by column chromotoghaphy over silica gel (eluent : CHCl₃). The desired fraction was collected and the eluent was evaporated. Yield: 34 g of 6-fluoro-3,4-dihydro-2*H*-1-benzopyran-2-methanol (interm.
- d) Reaction under N₂. To a stirred and cooled (-60°C; 2-propanone/CO₂ bath) mixture of ethanedioyl dichloride (0.1 mol) in CH₂Cl₂ (350 ml) was added sulfinylbis[methane] (30 ml) during 10 minutes. After stirring 10 minutes, a mixture of interm. 10 in CH₂Cl₂ (90 ml) was added during 5 minutes. After stirring for 15 minutes, N,N-diethylethanamine (125 ml) was added. When the mixture was warmed up to
- room temperature, it was poured out in water. The product was extracted with CH₂Cl₂. The organic layer was wased with water, HCl (1M), water, NaHCO₃ (10%) and water,

dried and evaporated. The residue was dissolved in diethyl ether, washed with water, dried, filtered and evaporated. The residue was purified by column chromotoghaphy over silica gel (eluent : CHCl₃). The desired fraction was collected and the eluent was evaporated. Yield: 21.6 g of 6-fluoro-3,4-dihydro-2*H*-1-benzopyran-2-carboxaldehyde (interm. 11)

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nButyllithium 1.6M (0.056 mol) was added slowly at -70°C to a solution of intermediate (5) (0.046 mol) in THF (100ml). The mixture was stirred at -70°C for 30 minutes. A suspension of interm. 11 (0.056 mol) in THF (100ml) was added slowly. The mixture was stirred at -70°C for 1 hour, then brought to room temperature, poured out into H_2O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (21g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 80/10; 15-35 μ m). The pure fractions were collected and the solvent was evaporated. Yield: 9.5g of interm. 12 (55%).

A mixture of intermediate (5) (0.1127 mol), 2-methoxyethanamine (0.2254 mol) and K_2CO_3 (0.2254 mol) in DMF (500ml) was stirred at 120°C for 15 hours and then cooled. The solvent was evaporated. The residue was taken up in CH_2Cl_2 and H2O. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated till dryness. The residue (33.53g) was purified by column chromatography over silica gel (eluent: CH_2Cl_2/CH_3OH 99.5/0.5; 15-40 μ m). Two fractions were collected and their solvents were evaporated. Yield: 5.7g of interm. 14 (38%) and interm. 13 (34%).

A mixture of intermediate (5) (0.0751 mol), thiomorpholine (0.0891 mol) and K₂CO₃ (0.15 mol) in DMF (200ml) was stirred at 120°C for 12 hours. The solvent was evaporated till dryness. The residue was taken up in CH₂Cl₂ and H₂O. The organic

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layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (26g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 80/20; 20-45 μ m). Two fractions were collected and their solvents were evaporated. The two fractions were combined. Yield: 9.4g of interm. 15 (37%); mp. 82°C.

Example A9

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a) 4-Aminobenzoic acid (0.219 mol) was added to a solution of sodium 3nitrobenzenesulfonate (0.118 mol) in H₂SO₄ 70% (230ml) and the mixture was stirred and refluxed. 2-propene-1,1-diol, 2-methyl-, diacetate (0.216 mol) was added dropwise and the mixture was refluxed for 4 hours. Ethanol (200ml) was added and the mixture was stirred at 80°C for 48 hours. The mixture was evaporated, the residue was poured into ice water/NH₄OH and extracted with CH₂Cl₂. The organic layer was dried (MgSO₄) and evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/2-propanol 99/1). The pure fractions were collected and evaporated. Yield: 21g of ethyl 3-methyl-6-quinolinecarboxylate (interm. 16) (45%). b) Interm. 16 (0.098 mol) in THF (270ml) was added at 0°C to a solution of LiAlH₄ (0.098 mol) in THF under N₂. When the addition was complete, water (10ml) was added. The precipitate was filtered off and washed with CH₂Cl₂. The organic layer was dried (MgSO₄), filtered off and evaporated. The product was used without further purification. Yield: 16.71g of 3-methyl-6-quinolinemethanol (interm. 17). c) MnO₂ (0.237 mol) was added to a solution of interm. 17 (0.096 mol) in CH₂Cl₂ (200ml) and the mixture was stirred at room temperature for 12 hours. The mixture was filtered through celite and the filtrate was stirred again with MnO₂ (20g) for 12 hours. MnO₂ (10g) was added again. The mixture was stirred for 12 hours. The mixture was filtered through celite and evaporated. The product was used without further purification. Yield: 11.71g of 3-methyl-6-quinolinecarboxaldehyde (71%) (interm. 18).

turnings (50ml) was added at 10°C to a mixture of THF (0.14 mol) in 1,1'-oxybisethane (10ml). A solution of interm. 18 (0.07 mol) in Mg turnings (100ml) was added carefully at 5°C, the mixture was poured into ice water and extracted with EtOAc. Yield: 11.34g of (±)-α-cyclohexyl-3-methyl-6-quinolinemethanol (63%) (interm. 19).

d) A solution of bromocyclohexyl (0.14 mol) in 1,1'-oxybisethane (50ml) and Mg

Example A10

Preparation of

A mixture of compound (5) (0.001507 mol), tributyl(1-ethoxyethenyl)stannane (0.00226 mol) and Pd(PPh₃)₄ (0.000151 mol) in 1,4-dioxane (5ml) was stirred at 80°C for 3 hours. Water was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. This product was used without further purification. Yield: 1.4g of interm. 20.

Example A11

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Preparation of

A mixture of compound (45) (prepared according to B6) (0.00125 mol) in NaOH 3N (5 ml) and iPrOH (1.7 ml) was stirred at room temperature overnight, then poured out into H_2O , acidified with HCl 3N and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was taken up in diethyl ether. The precipitate was filtered off and dried. Yielding: 0.26 g of intermediate 23 (56%). (mp.: 232°C)

15 Example A12

a. Preparation of

A mixture of 5-bromo-1H-indole-2,3-dione (0.221 mol) in NaOH 3N (500 ml0 was stirred at 80°C for 30 minutes, brought to room temperature and 2-pentanone (0.221 mol) was added. The mixture was stirred and refluxed for 1 hour and 30 minutes and acidified with AcOH until pH=5. The precipitate was filtered, washed with water and dried. Yielding 52.3 g of intermediate 24 and intermediate 25. (Total yielding: 80%).

nBuLi 1.6 M (0.0816 mol) was added dropwise at -78° C to a suspension of intermediate 25 (0.034 mol) and intermediate 26 (0.034 mol) in THF (300 ml) under N₂ flow. The mixture was stirred at -78° C for 30 minutes. nBuLi 1.6M (0.0816 mol) was added dropwise. The mixture was stirred for 1 hour. A mixture of intermediate 9 (0.102 mol) in THF (250 ml) was added slowly. The mixture was stirred for -78° C to -20° C, poured out into H₂O/HCl 3N and extracted with EtOAc. The organic layer was separated, dired (MgSO₄), filtered, and the solvent was evaporated till dryness. Yielding: 20.89 g of compound intermediate 26 and intermediate 27 (86%).

Example A13

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4-amino-3-methoxybenzoic acid (0.054 mol) was added portionwise at room temperature to a solution of 3-chloro-2-ethyl-2-butenal (0.065 mol) in AcOH (100ml). The mixture was stirred and refluxed for 8 hours and evaporated to dryness. The residue was taken up in CH₂Cl₂, water was added and the solution was basified by Et₃N. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue was crystallized from 2-propanone. The precipitate was filtered off and dried. Yielding: 2.5g of interm. 26 (18%).

CDI (0.012 mol) was added at room temperature to a solution of interm. 26 (0.011 mol) in CH_2Cl_2 (30ml). The mixture was stirred at room temperature for 1 hour. methoxyaminomethyl (0.012 mol) was added and the mixture was stirred at room temperature for 8 hours. H_2O was added. A precipitate was filtered off. The filtrate was extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered, and

the solvent was evaporated. The residue was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.95g of interm. 27 (31%) (mp.:148°C).

Example A14

Preparation of

Br (interm. 28)

4-Bromobenzenamine (0.034 mol) was added at room temperature to a solution of 3-chloride-2-ethyl-2-butanal (0.041 mol) in AcOH (60 ml). The mixture was stirred and refluxed for 8 hours, brought to room temperature and evaporated to dryness. The product was crystallized from EtOAc. The precipitate was filtered, washed with K2CO3 10% and taken up in CH2Cl2. The organic layer was separated, dried (MgSO4), filtered, and the solvent was evaporated. Yielding: 4,6 g of interm. 28 (54%).

Example A15

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a. Preparation of

A solution of KOH (0.326 mol) in H₂O (150ml) was added slowly at 5°C to a solution of 1,3-cyclohexanedione (0.268 mol) in H₂O (150ml). The temperature must not reach 12 °C. KI (2g) then 2-bromo-1-(4-nitrophenyl)ethanone (0.294 mol) were added portionwise. The mixture was stirred at room temperature for 48 hours. The precipitate was fitered, washed with H₂O then with diethyl ether and dried. Yielding: 63g (85%). A part of this fraction (1g) was crystallized from EtOH. The precipitate was filtered off and dried. Yielding: 0.5g of interm. 29 (42%) (mp.: 100°C).

b. Preparation of (interm. 30)

A mixture of interm. 29 (0.145 mol) in H₂SO₄ (40ml) was stirred at room temperature for 1 hour, poured out into ice, basified with NH₄OH and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue was crystallized from EtOH. The precipitate was filtered off and dried. Yielding: 31g (58%). A part of this fraction (1g) was crystallized from EtOH. The precipitate was filtered off and dried. Yielding: 0.7g of interm. 30 (58%) (mp.:200°C).

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A mixture of interm. 30 (0.039 mol), Raney Ni (10g) in EtOH (100ml) was hydrogenated at room temperature under a 3 bar pressure for 1 hour. The mixture was filtered over celite and washed with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (9.5g) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 4.6g (52%). The filtrate was evaporated. The residue (2.7g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH; 99/1; 15-40µm). Two fractions were collected and the solvent was evaporated. Yielding: 1.6g F1 and 1.2g F2. F2 was crystallized from EtOH. The precipitate was filtered off and dried. Yielding: 0.24g of interm. 31 (2%) (mp.:202°C).

Interm. 30 (0.02 mol) was added at room temperature to a solution of 3-chloro-2-ethyl-2-butenal (0.04 mol) in AcOH (50ml). The mixture was stirred and refluxed for 4 hours. The solvent was evaporated till dryness. The residue was crystallized from EtOAc. The precipitate was filtered off and dried. The residue was taken up in CH₂Cl₂. The mixture was basified with K₂CO₃ 10% and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue was crystallized from EtOH. The precipitate was filtered off and dried. Yielding: 2.5g of interm. 32 (40%).

Example A16

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A mixture of 2-(4-nitrophenyl)-1-phenylethanone (0.083 mol) and Raney Ni (20g) in EtOH (200ml) was hydrogenated at room temperature for 1 hour under a 3 bar pressure, then filtered over celite, washed with CH₂Cl₂/CH₃OH and dried. Yielding: 17.5g of interm. 33 (97%).

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B. Preparation of the final compounds

Example B1

Preparation of

(compound 306)

POCl₃ (0.024 mol) was added slowly at 5°C to DMF (0.024 mol). The mixture was stirred at room temperature for 30 minutes, then cooled to 5°C. 3-Oxo-butanoic acid ethyl ester (0.024 mol) was added slowly. The mixture was stirred at 5°C for 30 minutes. 1-(4-aminophenyl)-2-phenylethanone (0.024 mol) was added portionwise. The mixture was stirred at 90°C for 3 hours and dissolved in CH₂Cl₂. Ice water was added. The mixture was basified with NH₄OH and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue was crystallized from 2-propanone/diethyl ether. The precipitate was filtered off and dried. Yielding: 0.9 g of compound 306 (11%) (mp.:136°C).

Example B2

Preparation of

KMnO₄ (10g) was added portionwise at room temperature to a solution of

in tris(dioxa-3,6-heptyl)amine (1ml) and CH₂Cl₂ (100ml). The mixture was stirred at room temperature for 8 hours, filtered over celite, washed with CH₂Cl₂ and dried. The residue (6g, 100%) was crystallized from diethyl ether/petroleum ether. The precipitate was filtered off and dried. Yield: 2g of compound (2) (33%); mp. 82°C.

Example B3

a) Preparation of

nBuLi 1.6M (0.07 mol) was added slowly at -70°C to a solution of intermediate (5) (0.058 mol) in THF (150ml). The mixture was stirred at -70°C for 30 minutes. A solution of 2,3-dihydro-1*H*-Indene-2-carbonitrile (0.07 mol) in THF (100ml) was added slowly. The mixture was stirred at -70°C for 1 hour, brought slowly to room

temperature, hydrolized with H_2O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (22g) was purified by column chromatography over silica gel (eluent: CH_2Cl_2 / cyclohexane 80/20 to 100; 15-35 μ m). The pure fractions were collected and the solvent was evaporated. The second fraction was crystallized from 2-propanone/diethyl ether. The precipitate was filtered off and dried. Yield: 0.11g of compound (3). The filtrate was concentrated. Yield: 0.55g of compound (3); mp. 145°C.

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nBuLi 1.6M (0.022 mol) was added slowly at -70°C to a solution of intermediate (5) (0.018 mol) in THF (50ml). The mixture was stirred at -70°C for 1 hour, brought to -40°C, then cooled to -70°C. A solution of interm. 7 (0.018 mol) in THF (40ml) was added slowly. The mixture was stirred at -70°C for 1 hour, then brought to -20°C, hydrolyzed with H₂O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (6.5g) was purified by column chromatography over silica gel (eluent: toluene/EtOAc 90/10; 15-40µM). Two fractions (F1 and F2) were collected and the solvent was evaporated. F1 (2.4g) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 1.8g of compound (4) (29%); mp. 123°C. F2 (0.9g) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 0.2g of compound (5) (3%); mp. 120°C.

nBuLi 1.6M in exane (0.107 mol) was added dropwise at -78°C under N_2 flow to a mixture of intermediate (6) (0.089 mol) in THF. The mixture ws stirred at -78°C for 1 hour. A mixture of interm. 7 (150 ml) was added at -78°C under N_2 flow. The mixture was stirred at -78°C for 2 hours, brought to 0°C, poured out into H_2O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (31g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 20-45 μ m). Two pure fractions were collected and their solvents were evaporated. Yielding: 11 g of compound (7) (38%) and 8.2 g of compound (8) (28%).

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A solution of chloromethylbenzeen (0.0069 mol) in diethyl ether (8ml) was added slowly to a suspension of Mg (0.0069 mol) in a small amount of diethyl ether. The mixture was stirred at room temperature for 30 minutes (disparition of Mg), then cooled to 5°C. A solution of interm. 27 (0.0027 mol) in THF (8ml) was added slowly. The mixture was stirred at 5°C for 15 minutes, then at room temperature for 2 hours, poured out into H₂O and filtered over celite. The precipitate was washed with EtOAc. The filtrate was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (1g) was purified by column chromatography over kromasil (eluent: CH₂Cl₂ 100 to CH₂Cl₂/CH₃OH 99/1; 15-40µm). The pure fractions were collected and the solvent was evaporated. The residue (0.5g, 56%) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.14g of compound 503 (15%).

Example B4: examples of endgroup modifications

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A mixture of trans (compound 8)

(prepared according to example B3.c) (0.018 mol) in HCl 3N (60ml) and THF (60ml) was stirred at 60°C overnight. The mixture was basified with a K₂CO₃ 10% solution and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. Yield: 4.6g of compound (156) (82%).

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(prepared according to example B3.c) (0.0122 mol) in HCl 3N (40ml) and THF (40ml) was stirred and refluxed overnight, poured out into water, basified with K₂CO₃ 10% and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 40/60; 15-40μm). The pure fractions were collected and the solvent was evaporated. Yield: 2g of compound (9) (52%); mp. 226°C.

A mixture of compound (4) (0.0015 mol), 2-methoxyethanamine (0.003 mol) and K_2CO_3 (0.003 mol) in DMF (5ml) was stirred at 140°C for 48 hours. H_2O was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 60/40; 15-40 μ m). Two fractions were collected and the solvent was evaporated. Both fractions were crystallized separately from pentane. The precipitate was filtered off and dried. Yield: 0.05g of compound (10) (9%; mp. 115°C) and 0.057g of compound (11) (10%; mp. 107°C).

A mixture of compound (4) (0.0015 mol) in 2-(methylthio)ethanamine (2ml) was stirred at 120° C for 8 hours. K_2 CO₃ 10% was added. The mixture was extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (2.2g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 70/30; 15-40 μ m). Two fractions were collected and the solvent was evaporated. The first fraction was crystallized from diethyl ether/petroleum ether. The precipitate was filtered off and dried. Yield: 0.08g of compound (12) (14%); mp. 120°C. The second fraction was crystallized from diethyl

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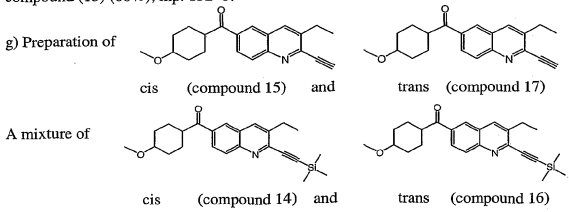
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ether. The precipitate was filtered off and dried. Yield: 0.18g of compound (13) (31%); mp. 125°C.

A mixture of compound (4) (0.001507 mol), ethynyltrimethylsilane (0.003013 mol), CuI (0.000151 mol) and Pd(PPh₃)₄ (0.000151 mol) in N,N-diethylethanamine (5ml) was stirred at 100° C for 24 hours. Water was added. The mixture was filtered over celite, washed with EtOAc and the filtrate was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.3g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue (0.3g) was crystallized from pentane. The precipitate was filtered off and dried. Yield: 0.11g of compound (14) (18%); mp. 114° C.

A mixture of compound (14) (0.013 mol) and KF (0.038 mol) in acetic acid (50ml) was stirred at room temperature for 2 hours. H_2O was added and the mixture was extracted with diethyl ether. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (4.4g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 70/30; 15-40 μ m). One fraction was collected and the solvent was evaporated. This fraction (3g, 73%) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 2.45g of compound (15) (60%); mp. 132°C.



prepared according to example B.7.a) (0.0056 mol) in KOH [1M, H₂O] (10ml) and methanol (30ml) was stirred at room temperature for 1 hour, poured out into water and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (2.2g) was purified by column

5 chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15 to 70/30; 15-40 μ m). Two fractions were collected and the solvent was evaporated.

The first fraction was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 0.2g of compound (15) (11%); mp. 133°C.

The second fraction was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 0.3g of compound (17) (16%); mp. 128°C.

h) Preparation of

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cis

A mixture of compound (4) (0.001205 mol), 2-propyn-1-ol (0.002411 mol), Pd(PPh₃)₄ (0.000121 mol) and CuI (0.000121 mol) in N,N-diethylethanamine (5ml) was stirred at 100°C for 2 hours. Water was added. The mixture was filtered over celite, washed with EtOAc and extracted aith EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.7g) was purified by column chromatography over silica gel (eluent: CH_2Cl_2/CH_3OH 98/2; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue was crystallized from petroleum ether and diethyl ether. The precipitate was filtered off and dried. Yield: 0.1g of compound (18) (23%); mp. 113°C.

i) Preparation of

solvents were evaporated.

A mixture of compound (4) (0.006027 mol) and KF (0.024108 mol) in DMSO (20ml) was stirred at 140°C. The solvent was evaporated till dryness. The residue was solidified in water and diethyl ether. The mixture was extracted with diethyl ether. The organic layer was separated, washed with diethyl ether, washed with a saturated solution of NaCl, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.7g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 15-40 μm). Three fractions were collected and their

The first fraction was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.21g of compound (19) (11%); mp. 92°C.

The second fraction was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.33g of compound (20) (17%); mp. 114°C.

A mixture of compound (4) (0.003013 mol), acetyl chloride (0.003315 mol) and sodium iodide (0.006027 mol) in CH₃CN (10ml) was stirred and refluxed for 1 hour. K₂CO₃ 10% was added. The mixture was extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1g) was purified by column chromatography over silica gel (eluent:

cyclohexane/EtOAc 80/20; 15-40 μ m). Two fractions were collected and their solvents were evaporated. The first fraction was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.12g of compound (21); mp. 110°C.

k) Preparation of

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A mixture of compound (21) (0.000898 mol), trimethylsilanecarbonitrile (0.001347 mol) and Pd(PPh₃)₄ (0.00009 mol) in N,N-diethylethanamine (5ml) was stirred at 100°C for 2 hours. Water was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄). filtered and the solvent was evaporated. The residue (0.4g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 80/20; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue (0.18g, 62%) was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.13g of compound (22) (45%); mp. 138°C.

cis

1) Preparation of

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A mixture of compound (4) (0.00603 mol), Pd(OAc)₂ (0.000603 mol), PPh₃ (0.00904 mol) and K₂CO₃ (0.012054 mol) in CO (gas) and methanol (40ml) was stirred at 90°C for 8 hours under a 5 bar pressure of CO. H₂O was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (6g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 100/0 to 98/2; 15-35μm). Four fractions (F1-F4) were collected and the solvent was evaporated. Yield: 0.13g (cis) F1; 0.02g F2 (cis, compound 25); 0.055g F3 (trans, 3%) and 0.11g F4 (trans; compound 26). F1 was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.03g of compound (23) (1%); mp. 91°C.

F3 was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.035g of compound (24) (1%); mp. 99°C.

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A mixture of compound (4) (0.009 mol) and Zn (0.027 mol) in acetic acid (30ml) was stirred at 60°C for 4 hours, filtered over celite, washed with CH_2Cl_2 , evaporated till dryness, solubilized in CH_2Cl_2 and washed with K_2CO_3 10%. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (4g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 75/25; 15-40 μ m). One fraction was collected and the solvent was evaporated. This fraction (1g 37%) was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: compound (25); mp. 88°C.

A mixture of compound (4) (0.001502 mol), Sn(CH₃)₄ (0.003004 mol) and Pd(PPh₃)₄ (0.00015 mol) in methylbenzene (5ml) was stirred and refluxed for 3 hours. K₂CO₃ 10% was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.7g)

was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; $15-40 \mu m$). Two fractions (F1 and F2) were collected and their solvents were evaporated. Yield: 0.27g (F 1, starting material) and 0.14g (F2). F2 was crystallized from pentane and petroleum ether. The precipitate was filtered off and dried. Yield: 0.08g of compound (27) (17%); mp. 110° C.

A mixture of compound (4) (0.001507 mol), tributylethenylstannane (0.002260 mol) and Pd(PPh₃)₄ (0.000151 mol) in dioxane (5ml) was stirred at 80°C for 8 hours. Water was added. The mixture was filtered over celite, washed with EtOAc and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.65g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 90/10; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.07g of compound (28) (14%); mp. 108°C.

A mixture of compound (5) (0.001507 mol), triphenyl(phenylmethyl)stannane (0.002260 mol) and Pd(PPh₃)₄ (0.000151 mol) in dioxane (5ml) was stirred at 80°C for 8 hours. Water was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.4g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/EtOAc 96/4; 15-40 μm). The pure fractions were collected and the solvent was evaporated. The residue (0.38g) was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.16g of compound (29) (28%); mp. 112°C.

cis

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A mixture of compound (4) (0.001507 mol), tributyl-2-thienylstannane (0.00226 mol) and Pd(PPh₃)₄ (0.0001507 mol) in dioxane (5ml) was stirred at 80°C for 8 hours. K_2CO_3 10% was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.7g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue (0.65g) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 0.35g of compound (30) (61%); mp. 142°C.

A mixture of compound (4) (0.0015 mol), 3-thienyl boronic acid (0.00226 mol), Pd(PPh₃)₄ (0.00015 mol) and dioxane was stirred and refluxed for 24 hours. K₂CO₃ 10% was added. The mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.8g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 80/20; 15-40μm). The pure fractions were collected and the solvent was evaporated. The residue (0.4g, 70%) was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.39g of compound (31) (68%); mp. 113°C.

cis

A mixture of compound (4) (0.003 mol), glycine methyl ester monohydrochloride (0.0066 mol) and Pd(PPh)₄ (0.0003 mol) in Et₃N (2ml) and toluene (10ml) was stirred at 100° C under 5 bar pressure of CO for 8 hours, filtered over celite, washed with CH₂Cl₂ and evaporated. The residue (2g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 80/20; 75-35 μ m). One fraction was collected and the solvent was evaporated. This fraction (1g 80%) was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.46g of compound (32) (37%).

A mixture of compound (4) (0.003 mol) and hydrazinecarboxaldehyde (0.0045 mol) in 1-butanol (15ml) was stirred and refluxed overnight, poured out into water and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/CH_3OH/NH_4OH$ 95/5/0.1; 15-40 μ m). Two fractions (F1 and F2) were collected and their solvents were evaporated. Yield: 0.3g F1 and 0.3g F2. F1 was crystallized from CH_3CN and diethyl ether. The precipitate was filtered off and dried. Yield: 0.102g of compound (33); mp. 224°C.

F2 was crystallized from CH₃CN and diethyl ether. The precipitate was filtered off and dried. Yield: 0.2g of compound (34); mp. 185°C.

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A mixture of compound 4 (0.015 mol) and NaN₃ (0.045 mol) in DMF (50ml) was stirred at 140°C for 2 hours. K_2CO_3 10% was added and the mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (6g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 60/40; 15-40 μ m). The first fraction was collected and the solvent was evaporated. The residue was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 1.26g of compound (35) (24%); mp. 160°C.

A mixture of compound (4) (0.009 mol) and thiourea (0.0099 mol) in ethyl alcohol (30ml) was stirred and refluxed for 12 hours and a solution of KOH (0.0149 mol) in H₂O (5ml) was added slowly. The mixture was stirred and refluxed for 1 hour, poured out into water and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (cyclohexane/EtOAc 70/30; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. Yielding: 1.1g of F1 (37%) and 0.4g of F2 (13%). F1 was crystallized from 2-propanone. The precipitate was

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filtered off and dried. Yielding: compound (36). F2 was crystallized from 2-propanone. The precipitate was filtered off and dried. Yielding: compound (37).

CH₃I (0.0034 mol) was added slowly at room temperature to a solution of compound (36) (0.0015 mol), compound (37) (0.0015 mol) and K₂CO₃ (0.0034 mol) in acetone (15ml). The mixture was stirred at room temperature for 8 hours. Water was added and the mixture was extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.2g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 15-40µm). The pure fractions were collected and the solvent was evaporated. Yielding: 0.6g F1 (57%), and 0.18g F2 (17%). F1 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.28g compound (38) (27%). F2 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.065g of compound (39) (6%).

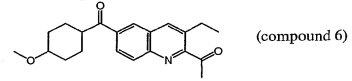
according to example B3b (0.0014 mol) in HCl 3N (5ml) and THF (5ml) was stirred and refluxed for a weekend, then poured out into H_2O , basified with K_2CO_3 and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. Yielding: 0.5g of F. This fraction F was crystallized from 2-propanone. The precipitate was filtered off and dried. Yielding: 0.35g of compound (40) (74%).

A mixture of compound (5) (0.045 mol), acetamide (0.90013 mol) and K_2CO_3 (0.225 mol) was stirred and refluxed at 200°C for 2 hours, cooled at room temperature, poured out into H_2O/CH_2Cl_2 ; and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated till dryness. The residue (14.4 g) was crystallized from CH_3OH . The precipitate was filtered off and dried. The filtrate was evaporated. The residue (11.27g) was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/CH_3OH/NH_4OH$ 96/4/0.1; 15-35 μ m). The pure fractions were collected and the solvent was evaporated. Yielding: 4.2 g of compound (188) (65%).

A mixture of compound (188) (0.00032 mol), benzoic acid (1.5 equiv., 0.00048 mol), 1-ethyl-3-(3'-dimethylaminopropyl)carbodiimide .HCl (1:1) (1.5 equiv., 0.00048 mol), N-hydroxybenzotriazole (1.5 equiv., 0.00048 mol) and Et₃N (1 equiv., 0.00032 mol) in CH₂CL₂ (2ml) was stirred at room temperature for 15 hours. The solvent was evaporated. The residue was purified by HPLC and the product fractions were collected and the solvent was evaporated. Yield: 0.066 g of compound (205) (49.50%).

aa) Preparation of

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trans

A mixture of interm. 20 (0.001507 mol) in HCl 3N (10ml) and THF (10ml) was stirred at room temperature for 8 hours, basified with K_2CO_3 10% and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.2g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/15; 15-40 μ m). The pure fractions were collected and the solvent was evaporated. The residue (0.4g) was crystallized from petroleum ether. The precipitate was filtered off and dried. Yield: 0.3g of compound (6) (58%); mp. 108°C.

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A mixture of compound 213 (prepared according to B4) (0.00305 mol) and CH₃ONa (30% in CH₃OH) (0.00916 mol) in CH₃OH (25ml) was stirred and refluxed for 15 hours then cooled to room temperature, poured out into H₂O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated till dryness. The residue (1.1g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc; 40/60; 15-40µm). Two fractions were collected and the solvent was evaporated. Yielding: 0.3g F1 and 0.5g F2 (50%) F2 was crystallized from diethyl ether/petroleum ether. The precipitate was filtered off and dried. Yielding: 0.26g F1 was crystallized from pentane. The precipitate was filtered off and dried. Yielding: 0.19g. This fraction was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH; 98/2; 15-40µm). The pure fractions were collected and the solvent was evaporated. Yielding: 0.11g. This fraction was purified by column chromatography over kromasil (eluent:CH₃OH/H₂O; 70/30). The pure fractions were collected and the solvent was evaporated. Yielding: 0.09g. (9%) This fraction was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.08g of compound 419 (8%).

Example B5

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Preparation of

cis (compound 42)

(trans) (compound 43)

Iodomethane (0.00456 mol) was added at 5°C to a mixture of compound (9) (0.0019 mol), compound (8) (0.0019 mol) and tBuOK (0.00456 mol) in THF (30ml) under N_2 flow. The mixture was stirred at room temperature overnight, poured out into H_2O and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 65/35; 15-40 μ m). Two fractions were collected and the solvent was evaporated. Yield: 0.35g of compound (42) (30%; mp. 125°C) and 0.35g of compound (43) (30%; mp. 116°C).

Example B6

a) Preparation of

cis (compound 44)

(trans) (compound 45)

NaH 60% (0.01068 mol) was added at 0° C under N_2 flow to a mixture of compound (8) and compound (9) (0.0089 mol). The mixture was stirred for 30 minutes.

Ethyl bromoacetate (0.01068 mol) was added at 0°C. The mixture was stirred at room temperature for 1 hour, hydrolized with water and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 60/40; 15-40 μ m). The desired fractions (F1-F4) were collected and the solvent was evaporated. Yield: 0.11g F1; 0.13g F2; 0.75g F3 and 0.8g F4.

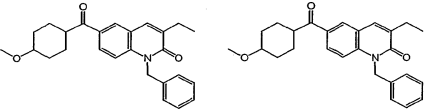
F3 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: compound (44); mp. 152°C.

F4 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: compound (45); mp. 147°C.

b) Preparation of

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cis (compound 46)

(trans) (compound 47)

Bromomethylbenzene (0.007 mol) was added dropwise at 0°C under N_2 flow to a solution of compound (8) and compound (9) (0.0064 mol) and NaH 60% (0.007 mol) in DMF (40ml). The mixture was stirred at room temperature for 1 hour, hydrolized with water and extracted with EtOAc. The organic layer was separated, washed with water, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 70/30; 15-40 μ m).

The desired fractions (F1-F4) were collected and the solvent was evaporated. Yield: 0.15g F1, 0.1g F2, 0.6g F3 (23%) and 0.8g F4.

F3 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yield: 0.13g of compound (46); mp. 137°C.

F4 was crystallized from DIPE and petroleum ether. The precipitate was filtered off and dried. Yield: compound (47); mp. 130°C.

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Example B7

a) 3-Chlorobenzenecarboperoxoic acid (0.088 mol) was added at 0°C to a solution of compound (48) (prepared according to example B2) (0.044 mol) in CH₂Cl₂ (200ml) and the mixture was stirred at room temperature for 12 hours. The mixture was washed with K₂CO₃ 10%. The organic layer was dried (MgSO₄), filtered off and evaporated. The residue was recrystallized from (C₂H₅)₂O. Yield: 8.2g of cyclohexyl(3-methyl-6-quinolinyl)methanone,1-oxide (compound 49) (69%). b) 4-Methyl benzenesulfonyl chloride (0.043 mol) was added to a solution of compound (49) (0.028 mol) in K₂CO₃ (400ml) and CH₂Cl₂ (400ml) and the mixture was stirred at room temperature for 1 hour. The mixture was extracted with CH₂Cl₂. The organic layer was dried (MgSO₄), filtered off and evaporated. The residue was recrystallized from (C₂H₅)₂O. Yield: 6.64g of 6-(cyclohexylcarbonyl)-3-methyl-2(1H)-quinolinone (compound 50) (85%); mp. 256.1°C.

Example B8

a) Preparation of

$$[1\alpha (A),4\alpha]$$
 (compound 51) $[1\alpha (B),4\alpha]$ (compound 52)

A mixture of compound (7) (0.0229 mol), hydroxylamine (0.0252 mol) and N,N-diethylethanamine (0.0252 mol) in ethanol (100ml) was stirred and refluxed for 6 hours, poured out into water and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was crystallized from CH_3CN . The precipitate was filtered off and dried. The residue was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/EtOAc$ 80/20; 15-40 μ m). Two fractions were collected and the solvent was evaporated. Yielding: 2.8g of compound (44) (36%; mp. 133°C) and 3g of compound (45) (38%; mp. 142°C).

b) Preparation of

$$(compound 53)$$

$$[1\alpha(Z),4\alpha]$$

Hydrazine (0.41 mol) was added at room temperature to a solution of compound (7) (0.015 mol) in ethanol (75ml). The mixture was stirred and refluxed for 1 night, poured out into water and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 98/2/0.1).

The pure fractions were collected and the solvent was evaporated. The residue was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.8g of compound (53) (15%); mp. 110°C.

Example B9

Preparation of

$$H_3CO$$
 (compound 520)

Procedure for compounds 400, 401, 402, 403, 404 and 405. A mixture of interm. 21 (prepared according to A11) (0.000269 mol), amantadine hydrochloride (0.000404 mol; 1.5 eq.), N'-(ethylcarbonimidoyl)-N,N-dimethyl-1,3-propanediamine hydrochloride (0.000404 mol; 1.5 equiv.), 1-hydroxy-1H-benzotriazole (0.000404 mol; 1.5 equiv.) and Et₃N (0.000269 mol) in CH₂Cl₃ (2 ml) was stirred at room temperature for 12 hours. The solvent was evaporated. The residue was purified by HPLC. The product fractions were collected and the solvent was evaporated. Yield: 0.063 g of compound 520 (46.37%).

Example B10

Preparation of

A mixture of intermediate 27 (0.0026 mol) and intermediate 26 (0.0026 mol) in EtOH (380 ml) and H₂SO₄ conc. (19 ml) was stirred and refluxed for 15 hours, the cooled to room temperature, poured out into ice water, basified with K₂CO₃ and extracted with EtOAc. The organic layer was separated, dried (MgSO4), filtered, and the solvent was evaporated. The residue (17.9 g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc; 80/20; 15-35μm). The pure fractions were collected and the solvent was evaporated. Yielding: 0.85 g of F1, 1.1 g F2 and 11.5 g of F3. F1 and F2 were crystallized separately from petroleum ether. The precipitate was filtered off and dried. Yielding: 0.34 g of compound 233.

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Example B11

Preparation of (compound 511)

A mixture of compound 22 (prepared according to B4) (0.004 mol) in HCl (3N) (20ml) and THF (20ml) was stirred and refluxed for 8 hours, poured out on ice, basified with NH₄OH and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (1.2g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH; 93/7/0.5; 15-40μm). Two fractions were collected and the solvent was evaporated. Yielding: 0.5g F1 (41%) and 0.4g of F2. F1 was crystallized from petroleum ether. The precipitate was filtered off and dried. Yielding: 0.17g of compound 511 (14%).

10 Example B12

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Preparation of (compound 514)

A mixture of compound 524 (prepared according to B9a) (0.0018 mol) and KOH 85% (0.0094 mol) in EtOH (15ml) was stirred and refluxed for 24 hours, poured out into $\rm H_2O$ and extracted with $\rm CH_2Cl_2$. The organic layer was separated, dried (MgSO4), filtered, and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: $\rm CH_2Cl_2/Cyclohexane~80/20;~15-40\mu m$). Two fractions were collected and the solvent was evaporated. Yielding: 0.35g F1 (64%) and 0.17g (SM) F1 was crystallized from diethyl ether. The precipitate was filtered off and dried. Yielding: 0.33g of compound 514 (60%) (mp.:185°C).

Example B13

Preparation of (compound 515)

A mixture of interm. 28 (0.019 mol), 2-benzofuranylboronic acid (0.028 mol), Pd(PPh₃)₄ (0.001 mol) and BHT (a few quantity) in dioxane (25ml) and Na₂CO₃ [2] (25ml) was stirred and refluxed for 8 hours and extracted with EtOAc. The aqueous layer was basified with NH₄OH and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (3.6g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 99/1;

 $15\text{-}40\mu\text{m}$). The pure fractions were collected and the solvent was evaporated. Yielding: 1.8g (33%). This fraction was crystallized from 2-propanone/diethyl ether. The precipitate was filtered off and dried. Yielding: 0.39g of compound 515 (7%) (mp.:134°C).

5 Example B14

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Preparation of (compound 526)

Triethylsilane (0.0012 mol) was added slowly at room temperature to a solution of interm. 32 (0.004 mol) in CF₃COOH (5ml) and AcOH (10ml). NaBH₄ (0.0012 mol) was added portionwise under N₂ flow. The mixture was stirred at room temperature for 8 hours, poured out on ice, basified with K_2CO_3 and extracted with CH_2Cl_2 . The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (1.2g) was purified by column chromatography over silica gel (eluent: CH_2Cl_2/CH_3OH 99/1; 15-40 μ m). Two fractions were collected and the solvent was evaporated. Yielding: 0.5g F1 (43%) and 0.4g F2. F1 was dissolved in iPrOH. HCl/iPrOH (1 eq) were added. The precipitate was filtered off and dried; Yielding: 0.32g of compound 526 (mp.: 248°C).

Example B15

Preparation of (compound 471)

A mixture of interm. 33 (0.082 mol) and 3-chloro-2-ethyl-2-butenal (0.098 mol) in AcOH (200ml) was stirred and refluxed for 8 hours. The solvent was evaporated till dryness. The residue was dissolved in CH_2Cl_2 and washed with K_2CO_3 10%. The organic layer was separated, dried (MgSO₄), filtered, and the solvent was evaporated. The residue (27g) was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/EtOAc$ 95/5 to 92/8; 15-35 μ m). Two fractions were collected and the solvent was evaporated. Yielding: 0.7g of F1 and 5.3g F2. F1 was crystallized from 2-propanone/diethyl ether. The precipitate was filtered off and dried. Yielding: 0.25g of compound 471 (2%) (mp.: 140°C).

Tables 1 to 8 list the compounds of formula (I-A) and (I-B) which were prepared according to one of the above examples.

Co.	Ex.	\mathbb{R}^2	$\hat{\mathbb{R}}^3$	R ⁴	\mathbb{R}^1	physical
l		K	IX.		T.	data
no.	no.				0	uata
54	B2	Cl	ethyl	Н	F. L. L.	_
3	ВЗа	Cl	ethyl	Н		mp. 145°C
55	B3b	Cl	ethyl	Н		mp. 131°C
56	B3b	Cl	ethyl	H		mp. 104°C
57	B3b	Cl	ethyl	Н	phenylethyl	mp. 100°C
58	B3b	Cl	ethyl	Н		mp. 126°C
59	B3b	Cl	ethyl	Н		mp. 150°C
60	B3b	Cl	ethyl	H		mp. 138°C
61	B3b	OCH₃	ethyl	Н		_
62	B3b	OCH ₃	ethyl	Н		mp. 130°C
63	B3b	OCH ₃	ethyl	н		mp. 116°С
64	B3b	Cl	ethyl	H	-(CH ₂) ₂ -O-CH ₃	mp. 82°C
65	B3b	OCH ₃	ethyl	H	1-methylcyclohexyl	mp. 82°C
66	B3b	OCH ₃	ethyl	H	3-methoxycyclohexyl	trans; mp. 94°C
67	B3b	OCH₃	ethyl	Н	3-methoxycyclohexyl	cis; mp. 108°C

Co.	Ex.	R ²	R ³	R ⁴	R ¹	physical
no.	no.			,		data
68	B3b	OCH ₃	ethyl	Н	4-(methylethoxy)-	(A), mp.
					cyclohexyl	82°C
69	B3b	OCH ₃	ethyl	H	4-[C(CH ₃) ₃]cyclohexyl	cis; mp. 92°C
70	B3b	OCH ₃	ethyl	H	4-[C(CH ₃) ₃]cyclohexyl	trans; mp. 108°C
71	ВЗЬ	OCH ₃	ethyl	Н	4-methylcyclohexyl	(B), mp. 92°C
72	B3b	OCH ₃	ethyl	Н	4-methylcyclohexyl	(A), mp. 80°C
2	В2	Cl	ethyl	Н	CH ₂ -CH(CH ₃) ₂	mp. 82°C
73	ВЗЬ	CI	ethyl	н	-CH ₂ -O-C ₂ H ₅	mp. 82°C
48	B2	H	methyl	Н	cyclohexyl	-
74	B4	I	ethyl	Н		_
75	B4	I	ethyl	Н		mp. 124°C
76	B4	I	ethyl	Н		mp. 138°C
77	B4	I	ethyl	Н	F	mp. 120°C
78	B4	CN	ethyl	н		mp. 128°C
79	B4	CN	ethyl	H	F	mp. 136°C
80	B4	CN	ethyl	H	C)	mp. 120°C
81	B4	CN	ethyl	н		mp. 139°C
82	B4	methyl	ethyl	Н		mp. 106°C
83	B4	methyl	ethyl	H		mp. 149°C
84	B4	methyl	ethyl	H	FU	mp. 118°C

Co.	Ex.	\mathbb{R}^2	R ³	R ⁴	R ¹	physical data
no. 85	no. B4	methyl	ethyl	Н	Ø.	mp. 180°C
86	В4	methyl	ethyl	H	phenylethyl	mp. 53°C
87	B4	methyl	ethyl	Н		mp. 87°C
88	B4	methyl	ethyl	Н	-CH ₂ -CH(CH ₃) ₂	mp. 68°C
89	B4	methyl	ethyl	Н		mp. 120°C
31	В4	3-thiazolyl	ethyl	H	4-methoxycyclohexyl	cis; 113°C
90	ВЗЬ	OCH ₃	H	Н	4-methoxycyclohexyl	trans, mp.
91	B3b	OCH ₃	H	H	4-methoxycyclohexyl	cis, mp. 100°C
92	B3b	OCH ₃	H	CH ₃	4-methoxycyclohexyl	cis; mp.
93	B3b	OCH ₃	Н	CH ₃	4-methoxycyclohexyl	trans; mp.
94	B3b	OCH ₃	methyl	H	4-methoxycyclohexyl	cis, mp. 96°C
95	ВЗЬ	OCH ₃	phenyl	Н	4-methoxycyclohexyl	cis; HCl (1:1), mp. 138°C
96	ВЗЬ	OCH ₃	propyl	Н	4-methoxycyclohexyl	trans; mp.
97	B3b	OCH ₃	propyl	Н	4-methoxycyclohexyl	
98	B3b	OCH ₃	methyl	Н	4-methoxycyclohexyl	cis; mp.
99	В4	N(CH ₃) ₂	ethyl	H	CH ₃	(B); mp. 102°C
100	B3b	Cl	ethyl	Н		mp. 114°C
101	B4	methyl	ethyl	H	4-butoxycyclohexyl	cis; mp. 86°C
102	ВЗь	Cl	ethyl	H		mp. 78°C

Co.	Ex.	R ²	R ³	R ⁴	\mathbb{R}^1	physical
no.	no.					data
103	ВЗЬ	Cl	ethyl	H		mp. 91°C
104	В4	N(CH ₃) ₂	ethyl	Н		mp. 103°C
105	В4	N(CH ₃) ₂	ethyl	Н	CL'ST	mp. 170°C
106	B3b	Cl	ethyl	Н		mp. 137°C
107	B3b	Cl	ethyl	H		mp. 137°C
108	В4	methyl	ethyl	ethyl	4-methoxycyclohexyl	cis; mp. 91°C
109	B4	methyl	ethyl	H	4-ethoxycyclohexyl	trans; mp.
110	В4	methyl	ethyl	H		mp. 90°C
111	B4	methyl	ethyl	Н		mp. 94°C
112	B4	methyl	ethyl	Н		mp. 176°С
113	B4	methyl	ethyl	H		mp. 106°C
114	B4	propyl	Н	Н	4-methoxycyclohexyl	cis; mp. 74°C
115	В4	methyl	ethyl	Н	4-ethoxycyclohexyl	cis; mp. 108°C
116	B4	methyl	ethyl	H		mp. 110°C
117	B3b	Cl	ethyl	Н		mp. 124°C
118	B3b	Cl	ethyl	Н		mp. 107°C
119	B3b	CI	ethyl	Н		mp. 129°C

Co.	Ex.	\mathbb{R}^2	R ³	R ⁴	\mathbb{R}^1	physical
no.	no.					data
120	B4	methyl	ethyl	Н		mp. 106°C
41	ВЗЬ	Cl	ethyl	H		trans; mp. 157°C
182	B3b	methyl	ethyl	H	но	cis; mp. 170°C
183	B3b	methyl	ethyl	Н	но	trans; mp. 144°C
184	ВЗЬ	methyl	ethyl	H	но	mp. 138°C
185	B3b	Cl	ethyl	H		mp. 120°C
186	ВЗЬ	Cl	ethyl	Н	> N	
187	B3b	methyl	ethyl	H	7	mp. 162°C
216	B4	CC≡N	ethyl	Н		mp.:160°C
217	B4	methyl	ethyl	Н		.ethanedioate (1:1); mp.:143°C
218	B4	I	ethyl	Н		mp.:102°C
219	B4	CC≡N	ethyl	Н	H ₃ C	mp.:115°C
220	B4	Cl	ethyl	Н	F	(A); mp.:107°C
221	B4	Cl	ethyl	Н	F	(B); mp.:113°C
222	B4	I	ethyl	Н	Û	mp.:206°C

Co.	Ex.	\mathbb{R}^2	\mathbb{R}^3	\mathbb{R}^4	\mathbb{R}^1	physical
no.	no.					data
223	B4	Cl	ethyl	Н	H ₃ C	(trans); mp.:117°C
224	B4	methyl	ethyl	Н	H ₃ C	(A); mp.:103°C
225	B2	Cl	ethyl	H		mp.:94°C
226	B3b	Cl	ethyl	H	C ₂ H ₅ O	(trans); mp.:157°C
227	ВЗс	methoxy	N	Н	H ₃ CO	mp.:204°C
228	B4	Cl	ethyl	H	H ₃ CO H ₃ CO	(A); mp.:136°C
229	B3b	n-propyl	Н	H	H ₃ CO H ₃ CO	(trans);.HCl (1:1); mp.:150°C
230	B3b	Cl	ethyl	Н	OCH ₃	mp.:116°C
231	B3b	Cl	ethyl	Н		mp.:120°C
232	B3b	CI	ethyl	Н		mp.:112°C
233	B10	<i>i</i> -propyl	1	C(=O)O- C ₂ H ₅	H ₃ CO H ₃ CO	(cis); mp.:91°C
234	B4	methyl	ethyl	Н		mp.:122°C
235	B4	methyl	ethyl	Н		mp.:106°C

Co.	Ex.	\mathbb{R}^2	\mathbb{R}^3	R ⁴	R ¹	physical
1 1			K	K	IX.	data
no.	no.		,1 1	TT		mp.:104°C
236	B4	methyl	ethyl	H		
					OCH ₃	
237	B4	methyl	ethyl	Н		mp.:90°C
257	D7	moury	Curyr	**		
						(cis);
238	В4	methyl	H	H		mp.:80°C
					H ₃ CO	
239	B3b	Cl	ethyl	H	H ₃ CO.	(trans);
239	D 30		Curyr	11		mp.:126°C
		The second secon			11.00	
240	ВЗЬ	Cl	ethyl	H	H ₃ CO Y	(cis); mp.:128°C
241	B4	methyl	ethyl	H	ÇH₃	(A); mp.:90°C
241	D4	linethy	Curyr	11		,
					H ₃ CO	
					ÇH ₃	(B);
242	B4	methyl	ethyl	H .		mp.:110°C
				}		
					H ₃ CO	mp.:134°C
243	B3b	Cl	ethyl	H		mp134 C
						·
					o´ o	,
		and the second s	and the second of the second o			mp.:127°C
244	B3b	Cl	ethyl	H		mp127 C
245	B4	NHC(=O)NH ₂	ethyl	H		(cis);
			,		<u> </u>	mp.:176°C
					H ₃ CO	(B)
246	B4	methyl	ethyl	H		(B)
					H ₃ C	
247	B3b	Cl	ethyl	Н		mp.:92°C
247	D 30		Curyi	11		
					0	
248	B4	methyl	ethyl	H		(A); mp.:80°C
1					H ₃ C	
-	Dei			TT	1130	(B);
249	B3b	Cl	ethyl	H		mp.:138°C
		1				

Co.	Ex.	\mathbb{R}^2	R ³	R ⁴	\mathbb{R}^1	physical
no.	no.					data
250	В4	methyl	ethyl	Н	O-npropyl	(trans); mp.:118°C
251	В4	methyl	ethyl	Н		(B);.HCl(1:1)
252	ВЗЬ	Cl	ethyl	Н	/propyl	(A)
253	B3b	Cl	ethyl	Н	ipropyl	(B)
254	B3b	methyl	ethyl	Н	(CH ₂) ₄	mp.:74°C
255	B4	methyl	ethyl	Н	O-npropyl	(cis); mp.:68°C
256	В4	methyl	ethyl	Н	но	mp.:210°C
257	В4	methyl	ethyl	Н	OCH ₃	mp.:113°C
258	B4	methyl	ethyl	Н	C ₂ H ₅	mp.:92°C
259	B3b	methyl	ethyl	Н	OCH ₃	mp.:115°C
260	B3b	methyl	ethyl	Н	OCH ₃	mp.:60°C
261	B3b	Cl	ethyl	Н	C ₂ H ₅	(A); mp.:86°C
262	ВЗЬ	Cl	ethyl	Н	C ₂ H ₅	(B); mp.:101°C

					2.02	
Co.	Ex.	R ²	R ³	R ⁴	\mathbb{R}^1	physical
no.	no.					data
263	ВЗЬ	methyl	ethyl	H	N(CH ₃) ₂	mp.:130°C
264	ВЗЬ	Cl	ethyl	H		(A); mp.:124°C
265	ВЗЪ	Cl	ethyl	H		(B); mp.:126°C
266	B4	N(CH ₃) ₂	ethyl	H	OCH ₃	(trans); mp.:102°C
267	В4	N(CH ₃) ₂	ethyl	Н	OCH ₃	(cis);.HCl(1:1); mp.:170°C
268	В4	methyl	ethyl	Н		(A);.HCl(1:1); mp.:206°C
269	B4	methyl	ethyl	H		mp.:104°C
270	B3b	methyl	ethyl	Н		mp.:117°C
271	В4	NHC2H5OCH3	ethyl	Н		
272	В4	methyl	ethyl	H	OCH3 N	-
273	B4	NH ₂	ethyl	Н		-
274	ВЗЪ	Cl	ethyl	Н	F	-
275	ВЗЬ	Cl	ethyl	Н	CF ₃	mp.:99°C

Co.	Ex.	R ²	R ³	R ⁴	R ¹	physical
no.	no.		ı			data
276	B3b	Cl	ethyl	Н	V	mp.:95°C
277	B4	methyl	ethyl	H	F	mp.:105°C
278	B3b	Cl	ethyl	H		mp.:141°C
279	В4	Cl	ethyl	Н	но	mp.:168°C
280	B4	Cl	ethyl	H	но	
281	В4	Cl	ethyl	Н	НО	mp.:140°C
282	B4	Cl	ethyl	Н		mp.:169°C
283	В4	methyl	ethyl	Н		mp.:96°C
284	B3b	Cl	CH ₂ N(CH ₃) ₂	Н		mp.:115°C
285	B4	methyl	ethyl	H	CH ₃	mp.:133°C
286	B4	methyl	CH₂OCH₃	Н	H ₃ CO	(trans); mp.:106°C
287	В4	methyl	CH ₂ N(CH ₃) ₂	H	H ₃ CO H ₃ CO	(cis); mp.:110°C
288	ВЗЬ	Cl	n-propyl	H		mp.:110°C

Co.	Ex.	R ²	R ³	R ⁴	R ¹	physical
no.	no.					data
289	B4	NH ₂	ethyl	Н		mp.:218°C
290	B4	methyl	n-propyl	H		mp.:90°C
291	ВЗъ	Cl	n-propyl	H	H ₃ CO H ₃ CO	(cis); mp.:128°C
292	B3b	Cl	n-propyl	H	H ₃ CO	(trans); mp.:104°C
293	B3b	Cl	ethyl	H		mp.:106°C
294	B4	methyl	n-propyl	Н	H ₃ CO	(cis); mp.:94°C
295	B4	methyl	CH ₂ N(CH ₃) ₂	Н		mp.:83°C
296	ВЗь	Cl	ethyl	H	S	mp.:99°C
297	ВЗЬ	Cl	ethyl	Н	T _s	mp.:110°C
298	B4	methyl	ethyl	Н	S	mp.:93°C
299	B4	methyl	ethyl	H	Š	mp.:105°C
300	B4	methyl	ethyl	H		mp.:114°C
301	ВЗь	methyl	ethyl	H		mp.:143°C
302	B4	methoxy	ethyl	H		mp.:93°C
303	B4	methyl	ethyl	H		mp.:82°C
304	B4	n-butyl	ethyl	H		-

Co.	Ex.	\mathbb{R}^2	\mathbb{R}^3	R ⁴	R ¹	physical
no.	no.					data
305	ВЗЪ	Cl	n-propyl	Н		mp.:125°C
306	B1	methyl	C(=O)OC₂H 5	Н		mp.:136°C
307	B4	methyl	n-propyl	H		mp.:81°C
308	B4	methoxy	n-propyl	Н		mp.:80°C
309	B4	I	n-propyl	H		mp.:120°C
310	B3d	methyl	ethyl	H	S	.HCl(1:1); mp.:129°C
311	B3b	Cl	H	Н		mp.:160°C
312	B3b	Cl	H	H	H ₃ CO H ₃ CO	(trans); mp.:145°C
313	B3b	Cl	H	H		mp.:103°C
314	B4	n-propyl	n-propyl	H		.HCl(1:1); mp.:150°C
315	B4	n-propyl	ethyl	Н		.HCl(1:1)
316	В4	n-propyl	H	Н		.HCl(1:1); mp.:140°C
317	ВЗЬ	Cl	Н	Н	S	mp.:168°C
318	В4	methyl	n-propyl	Н		.HCl(1:1); mp.:200°C
509	B3b	Cl	ethyl	Н		-
510	В4	methyl	ethyl	H		.H ₂ O(1:1)

	,	,				
Co.	Ex.	\mathbb{R}^2	\mathbb{R}^3	\mathbb{R}^4	\mathbb{R}^1	physical
no.	no.		-			data
513	В4	methyl	ethyl	H	H ₃ CO O	-
516	B4	Cl	ethyl	H	H ₃ C	mp.:120°C
517	B4	I	ethyl	H	CH ₂ CH(CH ₃) ₂	-
518	В4	Cl	ethyl	H	(F)	-
519	В4	Cl	ethyl	H	H ₃ CO CH ₃	(A+B)
521	B4	I	ethyl	Н		-
522	B4	methyl	ethyl	Н	J. I N	(A)
1	B4	methyl	ethyl	H	H ₃ CO CH ₃	(A)
525	B4	CI	ethyl	H	H ₃ CO O	
527	B4	F	ethyl	H		mp.: 116°C

Co.	Ex.	\mathbb{R}^2	x	physical data
no.	no.			
5	ВЗЬ	Cl	0	trans; mp. 120°C
121	B3b	1-piperidinyl	0	cis; HCl (1:1)
122	B3b	1-piperidinyl	0	trans; HCl (1:1); mp.
				128°C
123	B3b	4-thiomorpholinyl	0	cis; mp. 105°C
124	B3b	4-thiomorpholinyl	0	trans; mp. 115°C

Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
125	ВЗЬ	4-morpholinyl	0	trans; mp. 118°C
126	взь	4-morpholinyl	0	cis; mp. 118°C
127	B3b	-N(CH ₃) ₂	0	trans; mp. 96°C
128	ВЗЬ	-N(CH ₃) ₂	0	cis; mp. 114°C
4	B3b	Cl	0	cis; mp. 123°C
8	ВЗс	OCH ₃	0	trans, mp. 68°C
7	ВЗс	OCH₃	0	cis, mp. 116°C
6	B4	acetyl	0	trans; mp. 108°C
129	B4	acetyl	0	cis; mp. 106°C
11	B4	NH-(CH ₂) ₂ -OCH ₃	О	trans; mp. 107°C
10	B4	NH-(CH ₂) ₂ -OCH ₃	0	cis; mp. 115°C
12	B4	NH-(CH ₂) ₂ -SCH ₃	0	cis; mp. 120°C
13	B4	NH-(CH ₂) ₂ -SCH ₃	0	trans; mp. 125°C
14	B4	-C≡C-Si(CH ₃) ₃	0	cis; mp. 114°C
16	B4	-C≡C-Si(CH ₃) ₃	0	trans; mp. 108°C
15	B4	-C≡CH	0	cis; mp. 132-133°C
17	B4	-C≡CH	О	trans; mp. 128°C
18	B4	-C≡C-CH ₂ OH	O	cis; mp. 113°C
130	B4	-C≡C-CH ₂ OH	О	trans; mp. 108°C
19	B4	F	0	cis; mp. 92-99°C
20	B4	F	0	trans; mp. 114°C
21	B4	I	О	cis; mp. 110°C
22	B4	CN	0	cis; mp. 137-138°C
26	B4	H	О	trans
23	B4	-C(=O)-OCH ₃	0	cis; mp. 91°C
24	B4	-C(=O)-OCH ₃	0	trans; mp. 99°C
25	B4	H	0	cis; mp. 88°C
27	B4	methyl	0	cis; mp. 110-112°C
131	B4	methyl	О	trans; mp. 25°C
28	B4	ethenyl	O	cis; mp. 108°C
132	B4	ethenyl	О	trans; mp. 103°C
29	B4	phenyl	О	trans; mp. 112°C
30	B4	2-thienyl	0	cis; 142°C
133	B4	2-thiazolyl	0	cis; 108°C

Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
134	B4	2-furanyl	О	cis; mp. 105°C
51	B8a	OCH ₃	N-OH	$[1\alpha(A),4\alpha]$; mp. 133°C
52	B8a	OCH₃	N-OH	[1α(B),4α]; mp. 142°C
53	B8b	OCH₃	NNH ₂	$[1\alpha(Z),4\alpha]$; mp. 110°C
135	B4	NH ₂	О	cis; mp. 203°C
136	B4	NH ₂	0	trans; mp. 202°C
137	B4	-C(=O)-OCH(CH ₃) ₂	0	cis; mp. 105°C
138	B4	-C(=O)-OCH(CH ₃) ₂	0	trans; mp. 88°C
38	B4	SCH ₃	0	cis; mp. 124°C
39	B4	SCH₃	0	trans; mp. 116°C
32	B4		О	cis; mp. 130°C
		Ö		10000
139	B4	ethyl	0	cis; mp. 180°C
188	B4	NH ₂	0	cis + trans
189	B4	OCH ₃	O	cis; mp. 154°C
190	В4	OCH3	О	trans; mp. 156°C
191	B4	NH ONH	0	cis; mp. >260°C
192	B4	NH ONH	О	.H2O (1:1); trans; mp. 248°C
193	В4	H CH₃	O	cis; mp. 224°C
194	B4	₩ CH ₃	0	trans; mp. 234°C
195	B4	H OC₂H₅	О	cis; mp. 108°C

Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
196	B4	H N OC ₂ H ₅	0	trans; mp. 127°C
197	B4	√ly s \ \	О	cis; mp. 150°C
198	B4	√H y s \ \	О	trans; mp. 90°C
199	B4		0	LC/MS [M+H] ⁺ ; 475.4
200	B4	TH S N	О	LC/MS [M+H] ⁺ ; 464.3
201	B4		О	LC/MS [M+H] ⁺ ; 523.3
202	B4	H s	О	LC/MS [M+H] ⁺ ; 465.3
203	B4	$ OC_2H_5$	О	LC/MS [M+H] ⁺ ; 475.4
204	B4	H S N	О	LC/MS [M+H] ⁺ ; 465.3
205	B4		О	-
319	B4		О	(cis);.ethanedioate(1:1); mp.:160°C
320	В4		О	(cis); mp.:150°C
321	B4	methoxy	CH ₂	(cis);.HCl(1:1); mp.:118°C
322	B4	n-butyl	О	(cis);.HCl(1:1); mp.:158°C

Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
323	B4	OCH ₃	0	-
324	B4		О	-
325	B4	CH ₃	0	-
326	B4	H N O CH₃	0	-
327	B4	N → CI	О	-
328	B4	_H	0	-
329	B4	N S S S S S S S S S S S S S S S S S S S	0	-
330	B4	N(CH ₃) ₂	О	
331	В4	NH CO	О	-
332	B4	H _S s	О	•
333	B4	N CH₃	0	-
334	B4	H C	0	_
335	B4	H CH ₃	О	-

Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
336	B4	The state of the s	О	-
337	B4	H N	О	-
338	B4	CH ₃	0	-
339	B4	, N	О	-
340	B4	H ₃ C H N	О	-
341	B4	OCH ₃	O	-
342	B4		О	_
343	B4	N OCH₃	0	-
344	B4		О	-
345	B4	CH ₃	О	_
346	B4		О	-
347	B4	CF ₃	О	-
348	B4	CH ₂ OC(=O)CH ₃	О	(cis); mp.:74°C
349	B4	H O CH ₃	0	-

_		-2	47	
Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			
350	B4	H N O CH₃	О	-
351	B4	CH ₃ O N O CH ₃	О	-
352	B4	H. N	0	-
353	B4	CH ₃	О	(A);.HCl(1:2).H2O(1:1); mp.:166°C
354	B4	CH ₃	О	(cis)
355	B4	H CH ₃	0	-
356	B4		0	-
357	В4	CH ₃	O	-
358	В4	- N	Ο.	-
359	B4		О	
360	B4		0	.= '
361	B4	N(CH ₃) ₂	O	-
362	B4	CH ₃	0	-
363	B4	H H CH ₃	0	-

Co.	Ex.	\mathbb{R}^2	X	physical data
nọ.	no.			
364	B4	CH ₃ CH ₃	О	_
365	B4	H H O CH ₃	O	_
366	B4	_H CH₃	О	-
367	B4	H H CH ₂	О	-
368	B4		O	-
369	B4	N(CH ₃) ₂	О	-
370	B4	H H O CH ₃	О	
371	B4		О	-
372	В4	H H CH3	О	-
373	B4	H H S CH ₃	О	-
374	B4	H H O CH3	О	-
375	B4	O S S CI	0	-

	-		I	T
Co.	Ex.	\mathbb{R}^2	X	physical data
no.	no.			, .
376	B4	0=s N H O	О	-
377	B4 ,	CF ₃	О	-
378	B4	OH S CH ₃	0	_
379	B4	N N N N N N N N N N N N N N N N N N N	О	-
380	B4	CH ₃	О	-
381	B4	OHO CH3	О	-
382	В4	OH3 CH3	О	-
383	В4	OCH ₃	0	(cis); mp.:148°C
384	В4	OCH3	О	(trans); mp.:141°C
385	В4		0	mp.:130°C
386	B4	S _{CH₃}	O	(cis); mp.:140°C
387	B4	-N	О	(trans); mp.:155°C

Co.	Ex.	Y.	\mathbb{R}^1	physical data
no.	no.			
140	B4	О		mp. 220°C
141	B4	О		mp. 213°C
142	B4	О		mp. 148°C
143	B4	0	1-methylcyclohexyl	mp. 195-210°C
144	B4	0	3-methoxycyclohexyl	cis; mp. 156°C
145	В4	0	3-methoxycyclohexyl	trans; mp. 156-163°C
146	B4	0	4-(dimethylethyl)cyclohexyl	mp. 230°C
147	B4	0	4-(methylethoxy)cyclohexyl	mp. 186°C
148	В4	0	4-methylcyclohexyl	trans; mp. 214°C
36	B4	S	4-methoxycyclohexyl	cis; mp. 224°C
37	B4	S	4-methoxycyclohexyl	trans; mp. 220°C
149	B4	O		mp. 188°C
40	B4	О		mp. 192°C
150	B4	О		cis; mp. 226°C
151	B4	О		trans; mp. 226°C
152	B4	О		mp. 213°C
153	B4	О		mp. 200°C

Co.	Ex.	Y.	\mathbb{R}^1	physical data
no.	no.			ř ,
154	В4	О		mp. 210°C
155	B4	0	4,4-dimethylcyclohexyl	mp. 242°C
388	B4	О	CH ₂ CH(CH ₃) ₂	mp. 189°C
389	B4	О		mp. 228°C
390	B4	O		mp. 197°C
391	B4	О		mp. 145°C
392	B4	О	F	mp. 192°C
393	B4	О	H ₃ C _O F	(B); mp.:224°C
394	B4	О	H ₃ C O	(A); mp.:201°C
395	B4	О	H ₃ C CH ₃	(A); mp.:207°C
396	B4	О		mp.:212°C
397	B4	О		(B); mp.:238°C
398	B4	О		mp.:234°C
399	B4	О	CH ₃	(cis); mp.:192°C

$$\mathbb{R}^{4}$$

$$\mathbb{R}^{5}$$

Co.	Ex.	\mathbb{R}^3	R ⁴	\mathbb{R}^5	R	physical data
no.	no.					
156	B4	ethyl	Н	H	OCH ₃	trans; mp. 252°C
157	В4	Н	н	H	OCH ₃	(cis + trans);
						mp. 244°C
158	B4	Н	methyl	H	OCH ₃	cis; mp. >260°C
159	B4	methyl	Н	H	OCH ₃	cis; mp. 254°C
160	B4	methyl	Н	H	OCH ₃	trans; mp. >260°C
161	B4	propyl	H	H	OCH ₃	mp. 208°C
162	B4	propyl	Н	H	OCH ₃	trans; mp. 232°C
9	B4	ethyl	Н	H	OCH₃	cis; mp. 224-226°C
43	B5	ethyl	H	CH₃	OCH₃	trans; mp. 116°C
42	B5	ethyl	H	CH₃	OCH ₃	cis; mp. 125°C
44	В6	ethyl	H	CH ₂ -COOC ₂ H5	OCH ₃	cis; mp. 152°C
45	B4	ethyl	H	CH ₂ -COOC ₂ H5	OCH ₃	trans; mp. 147°C
46	B4	ethyl	H	benzyl	OCH ₃	cis; mp. 137°C
47	B4	ethyl	H	benzyl	OCH ₃	trans; mp. 130°C
50	B7	methyl	Н	Н	H	mp. 256.1°C
163	B4	ethyl	ethyl	H	OCH ₃	cis; mp. 221°C
164	B4	ethyl	ethyl	H	OCH ₃	cis; mp. 221°C
165	B4	ethyl	ethyl	H	OCH ₃	trans; mp. 215°C
166	R4	ethyl	H	L N	OCH ₃	LC/MS [M+H] ⁺ ;
100		Cary			00113	429.4
167	B4	ethyl	H	H 9	OCT	LC/MS [M+H] ⁺ ;
	}			Ĭ, N	OCH₃	451.3
168	B4	H	H	H	OCH₃	cis; mp. 106°C
169	R1	ethyl	H	H		LC/MS [M+H] ⁺ ;
109	1 24	Curyi	**		OCH₃	409.3
<u></u>	<u> </u>			0		

Co.	Ex.	\mathbb{R}^3	R ⁴	\mathbb{R}^5	R	physical data
no.	no.					
400	В9	ethyl	Н	TH.	OCH₃	-
401	В9	ethyl	Н		OCH ₃	-
402	В9	ethyl	Н		OCH ₃	-
403	В9	ethyl	H		OCH ₃	-
404	В9	ethyl	Н		OCH ₃	-
405	В9	ethyl	H	CH ₃	OCH ₃	\ -
406	B4	ethyl	Н		OCH ₃	-
407	В4	ethyl	H	T S S	OCH ₃	
408	B4	ethyl	H		OCH ₃	-
409	ВЗЪ	(CH ₂) ₃	Н	H	OCH ₃	mp.:168°C
410	B4	CH₂OCH₃	Н	H	OCH ₃	mp.:194°C
508	B4	ethyl	H	H OC ₂ H ₅	OCH₃	_
520	B9	ethyl	н	S S	OCH₃	

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Co.	Ex.	R ⁴	R ¹ .	X	physical data
no.	no.) 	<u>u</u>
33	B4	H	methoxycyclohexyl	СН	cis; mp. 224°C
34	B4	H	methoxycyclohexyl	СН	trans; mp. 185°C
35	B4	H	methoxycyclohexyl	N	cis; mp. 160-172°C
170	B4	H	methoxycyclohexyl	N	trans; mp. 146°C
171	B4	Н		N	(B); mp. 165°C
172	B4	H	methylcyclohexyl	N	cis+trans; mp. 143°C
173	B4	ethyl	methoxycyclohexyl	N	cis; mp.:126°C
411	B4	H		N	mp.:109°C
412	B4	H		N	mp.:180°C
413	В4	Н		N	(A)
414	B4	H	H ₃ CO	N	mp.:156°C

ŔÌ	_			
Co.	Ex.	R	L	physical data
no. 49	no.	Н	CH ₃	
174	ВЗЬ	OCH ₃		cis; mp.115°C
175	ВЗЬ	OCH ₃		trans; mp.141°C
176	B3b	OCH ₃		cis; mp.149°C
177	B3b	OCH ₃		mp.126°C
178	ВЗЬ	OCH ₃	TINS .	trans; mp.160°C
179	B3b	OCH₃	J CI	cis; mp.119°C
180	B3b	OCH ₃	TO.	trans; mp.124°C
181	B3b	OCH ₃	III.	trans; mp.92°C
206	B3b	ОСН₃		cis; m.p. 144°C

Co.	Ex.	R	L	physical data
no.	no.			
207	B3b	OCH ₃	CH ₃	trans; m.p. 125°C
208	ВЗЪ	OCH ₃	CH ₃	cis; m.p. 127°C
209	B3b	OCH ₃		cis; m.p. 101°C
210	B3b	OCH₃	H ₃ C N CI	cis; m.p. 104°C
211	B3b	OCH ₃	H ₃ C N CI	trans; m.p. 134°C
212	B4	OCH₃		cis; m.p. 141°C
213	B4	OCH ₃	H ₃ C	trans; m.p. 215°C
214	B4	OCH₃	H ₃ C N	cis; m.p. 139°C
215	B3b	OCH ₃	J N CI	trans
415	B3b	OCH ₃		(cis); mp.:136°C
416	B3b	OCH ₃	TINS	(cis)

Co.	Ex.	R	L	physical data
no.	no.			
417	В4	OCH₃	CH ₃	(cis); mp.:149°C
418	B3b	OCH ₃	TIN'S	(trans); mp.:132°C
419	В4	OCH ₃	H ₃ C N O CH ₃	(cis); mp.:217°C
420	В3ь	OCH ₃		(cis);.HCl(1:1); mp.:200°C
421	B4	ОН	TIN'S	(cis); mp.:215°C
422	B4	ОН	TINS	(trans); mp.:178°C
423	B3b	OCH₃	N CH ₃	mp.:160°C
424	B3b	OCH ₃		(cis); mp.:106°C
425	B3b	OCH ₃		(trans); mp.:120°C
426	B3b	ОСН₃		(cis); mp.:121°C
427	B3b	Н	TINS	mp.:156°C
428	B3b	ОСН₃		(cis); mp.:156°C
429	B3b	OCH ₃		(trans); mp.:197°C

Co.	Ex.	R	L	physical data
no.	no.			
430	ВЗЪ	CH ₃	TINIS	(B)
431	ВЗЬ	CH ₃	T S	(A)



Co.	Ex.	R ¹	L	physical data
no.	no.			
432	B4			mp.:128°C
433	B4	8		mp.:175°C
434	B4			mp.:170°C
435	B4			mp.:103°C
436	B4			mp.:151°C
437	B4	OCH ₃		(trans); mp.:110°C
438	B4			mp.:150°C
439	B4		N S	mp.:150°C
440	B4	OCH ₃		(cis)
441	B4		T S	mp.:166°C

				
Co.	Ex.	\mathbb{R}^1	L	physical data
no.	no.			
442	B4	N(CH ₃) ₂		mp.:173°C
443	B4			mp.:208°C
444	B4	CH ₂		mp.:149°C
445	B4			mp.:133°C
446	ВЗЬ			mp.:150°C
447	B3b		N S	mp.:165°C
448	B3b		T S	mp.:147°C
449	B3b		T S	mp.:154°C
450	B3b		T S	mp.:157°C
451	B4	H ₃ C N	T N S	mp.:190°C
452	B4			mp.:187°C
453	B3b	Br	T S	mp.:200°C
454	B3b			mp.:160°C

Co.	Ex.	R ¹	L	physical data
no.	no.	,		
455	ВЗЪ		NO CH3	mp.:139°C
456	B3b		T No	(A); mp.:174°C
457	B3b			(B); mp.:160°C
458	ВЗЪ		N CH ₃	mp.:184°C
459	В4	NC NC	C(CH ₃) ₃	
460	B4	H ₃ CO 0		mp.:134°C
461	B4			(B); mp.:156°C
462	B4		T S	mp.:153°C
463	B3b			mp.:161°C
464	B4	S	T S	mp.:135°C
465	B4	S		mp.:131°C
466	B3b		T S	.HCl(1:1); mp.:206°C
467	B3d		O C(CH ₃) ₃	mp.:142°C
468	B4			.hydrate(1:1); mp.:104°C
469	ВЗъ	dimethylethyl	T S	mp.:104°C

Co.	Ex.	R ¹	L	physical data
no.	no.	, , , , , , , , , , , , , , , , , , , ,		
470	B3b		T S	mp.:161°C
472	ВЗЬ		N	mp.:144°C
473	B4			mp.:143°C
474	B4	F		mp.:196°C
475	B4	F		mp.:162°C
476	B4	CH ₃	N	mp.:171°C
477	B4 ⁻	F		mp.:155°C
478	B2	trimethylmethyl	T S	mp.:124°C
479	B4	H ₃ CO F	T S	(A); mp.:146°C
480	B4	H ₃ CO F	T S	(B); mp.:162°C
481	B4	CH ₃		(A); mp.:129°C
482	B4			mp.:115°C
483	B2	F	T S	mp.:187°C
484	B2	F	T S	mp.:162°C
485	B4	H ₃ CO CH ₃	T S	(A); mp.:130°C

Co.	Ex.	\mathbb{R}^1	L	physical data
no.	no.		-	
486	B4	H ₃ CO CH ₃		(A); mp.:124°C
487	B4	H ₃ CO CH ₃		(B); mp.:128°C
488	B4	H ₃ CO F		mp.:85°C
489	B2	H ₃ C S		mp.:150°C
490	B4	H ₃ CO		(A); mp.:117°C
491	В2			mp.:220°C
492	B4	CH₃	TINIS	mp.:136°C
493	B2	N(CH ₃) ₂		mp.:131°C
494	B4	CH₃		(A); mp.:125°C
495	B4		TINIS	mp.:135°C
496	B4		T S	mp.:139°C
497	В4			mp.:127°C
498	B4	Br		mp.:195°C
499	B2			mp.:201°C

Co.	Ex.	R ¹	L	physical data
no.	no.			_
500	B3b		N O CH ₃	mp.:143°C
501	B3b			mp.:137°C
502	B2			mp.:210°C
503	B3d		CH ₃ OCH ₃	mp.:134°C
504	B2		H ₃ C N O	mp.:163°C
505	В4		NH	mp.:142°C
506	B2		H ₃ C N S	mp.:139°C
507	B4		T S	mp.:171°C
512	B3b		CLNS	-
523	B3b	F	T S	-

Table 8:

Co.	Ex.	Structure	physical data
no.	no.		
511	B11	CH ₃	

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Co.	Ex.	Structure	physical data
no.	no.		<u></u>
514	B12		-
515	B13	CH ₃	-
524	B9a	F OH	mp.:185°C
471	B15	CH ₃	(E)
526	B14	CH ₃	.HCl(1:1)

C. Pharmacological example

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15

Signal transduction at the cloned rat mGluR1 receptor in CHO cells

CHO cells expressing the mGluR1 receptor were plated in precoated black 96-well plates. The next day, the effect of the present compounds on glutamate-activated intracellular Ca²⁺ increase was evaluated in a fluorescent based assay. The cells were loaded with Fluo-3 AM, plates were incubated for 1 hour at room temperature in the dark, cells were washed and the present compounds were added onto the cells for 20 minutes. After this incubation time, the glutamate-induced Ca²⁺ rise was recorded for each well in function of time using the Fluorescent Image Plate Reader (FLIPR, Molecular Devices Inc.). Relative fluorescence units were recorded and average data graphs of quadruple wells were obtained. Concentration-response curves were constructed based on peak fluorescence (maximum signal between 1 and 90 secondes) for each concentration of tested compound. pIC₅₀ values are the –log values of the concentration of the tested compounds resulting in 50% inhibition of the glutamate-induced intracellular Ca²⁺ rise.

The compounds according to the present invention exhibited a pIC₅₀ value of at least 5.

The compounds that are included in the Tables 1-8 exhibited a pIC₅₀ value of at least 6.

A particular group of compounds exhibited a pIC_{50} value between 7 and 8. It concerns the compounds listed in Table 9.

Table 9:

	TC
Com.nr.	pIC ₅₀
463	7.98
441	7.95
334	7.95
22	7.94
421	7.94
15	7.93
440	7.93
139	7.93
178	7.92
338	7.91
87	7.90
462	7.90
394	7.90
423	7.89
21	7.87
220	7.87
479	7.86
483	7.86
485	7.84
9	7.84
110	7.84
248	7.84
341	7.83
163	7.81
433	7.79
238	7.79
224	7.78
437	7.78
498	7.78
449	7.77
242	7.76
346	7.74
182	7.73
486	7.73
447	7.72
7	7.72
175	7.71
475	7.71

Com.nr.	pIC ₅₀
281	7.63
487	7.63
299	7.63
431	7.61
98	7.57
464	7.57
446	7.56
251	7.55
484	7.54
494	7.53
128	7.52
344	7.52
161	7.49
298	7.48
454	7.45
456	7.45
277	7.44
91	7.43
356	7.42
229	7.41
333	7.41
326	7.41
369	7.40
430	7.39
435	7.38
35	7.36
228	7.36
429	7.36
117	7.35
291	7.35
313	7.35
280	7.34
460	7.34
482	7.34
343	7.33
343 425	7.33 7.32
473	7.32
473 287	7.31

Com.nr.	pIC ₅₀
89	7.25
108	7.25
373	7.25
255	7.23
527	7.23
303	7.22
296	7.22
221	7.21
193	7.21
14	7.20
131	7.19
438	7.19
148	7.18
496	7.18
236	7.17
332	7.17
481	7.16
191	7.16
457	7.14
20	7.14
145	7.13
268	7.13
512	7.13
474	7.13
10	7.11
307	7.11
426	7.11
466	7.10
97	7.08
83	7.08
434	7.08
300	7.08
199	7.07
290	7.06
112	7.05
348	7.05
286	7.05 7.03
442	7.03

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Com.nr.	pIC ₅₀
480	7.71
213	7.70
239	7.70
241	7.67
461	7.65
115	7.64
445	7.63

Com.nr.	pIC ₅₀
448	7.31
243	7.29
323	7.28
159	7.28
289	7.27
184	7.26
436	7.26

Com.nr.	pIC ₅₀
422	7.02
283	7.02
318	7.02
36	7.00
396	7.00

A particular group of compounds exhibited a pIC_{50} value of at least 8. It concern the compounds listed in Table 10.

<u>Table 10</u>:

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Comp.	Structure	pIC50
416	(CIS)	8.587
27	(CIS)	8.527
174	(CIS)	8.49
506	O O O O O O O O O O O O O O O O O O O	8.48

Comp.	Structure	pIC50
nr. 25		8.45
4	(CIS)	8.4
19	(CIS)	8.38
429	(CIS)	8.38
424	(CIS)	8.355
176	(CIS)	8.33

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210 210 210 8.315 114 8.28 488 8.27 504 60 8.27 477 8.25 432 60 60 8.237	Comp.	Structure	pIC50
114 (CIS) 488 504 60 8.27 477 60 8.25 432 60 8.237 8.237			8.315
114 (CIS) 488 8.27 504 (CIS) 8.27 477 (CIS) 8.27 8.27 432 (CIS) 8.27 8.27 8.27		(CIS)	
488 8.27 504 8.27 477 8.25 432 8.237	114		8.28
477 432 0 8.25 8.237	488	0	
432 8.237 214 8.233	504	O O O O O O O O O O O O O O O O O O O	8.27
214 8.233	477	o F	8.25
	432	O CONTRACTOR OF THE PARTY OF TH	
(CIS)	214		8.233

Comp.	Structure	pIC50
nr. 465		8.145
403	0	0.143
	s	
	, NO.	
135	O II	8.14
	N NH ₂	
,	(CIS)	
420	0	8.135
	N N	
	(CIS) Hydrochloride (1:1)	
292		8.13
	0	
	N N	
	(CIS)	,
427	(00)	8.115
	o 11	
		,
200		8.005
208	0	8.095
	N N N	
	l l	
	(CIS)	

Comp.	Structure	pIC50
419	(CIS)	8.065
455	(CIS)	8.055
418	(TRANS)	8.045
497		8.025
439	O O O O O O O O O O O O O O O O O O O	8.023
237	C	8.01

Comp.	Structure	pIC50
nr.		
499		8
	o F	

Cold allodynia test in rats with a Bennett ligation.

Surgery:

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Male SD rats, weighing 240 - 280 g at the time of surgery were used.

For surgery, the animals were anaesthetised with Thalamonal (1 ml; subcutane) and sodium pentobarbital (40 mg/kg; intraperitoneal (IP)). The common sciatic nerve of the left hindpaw was exposed at the level of the middle of the thigh by blunt dissection through the biceps femoris. Proximal to the sciatic's trifurcation, about 7 mm of nerve was freed and four loose ligatures with 4.0 chromic gut were placed around the sciatic nerve. Great care was taken to tie the ligatures such that the diameter of the nerve was barely constricted. After surgery, the animals received 1.25 mg/kg naloxone IP.

Cold plate testing:

Cold plate testing was performed on a metal plate of 30 X 30 cm with transparent acrylic walls around it. The cold plate was cooled to 0.0 (± 0.5) °C using a Julabo F25 cooler. For testing, the animal was placed on the cold plate and the duration of lifting of both the left and the right hindpaw was measured during 5 minutes. The difference in lifting time between the ligated and non-ligated paw was calculated.

Testing procedure:

At least one week after the operation, animals were placed on the cold plate test and a pre-drug measurement was taken. Animals having a difference in lifting time > 25 secondes between the ligated and the non-ligated paw were selected for drug testing. These selected animals were injected IP with a compound of the present invention and were retested after 60 minutes (post drug test). The results obtained during the post drug test were expressed as a percentage of those of the predrug test.

The data were analysed in terms of all or none criterion (based on the results of control animals) with the limits being:

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Inhibition: (post-drug/pre-drug)*100 < 40 %

Antagonism: (post-drug/pre-drug)*100 < 25 %

Compound (27) showed antagonism at a dose of 2.5 mg/kg bodyweight.

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Claims

1. A compound of formula

$$R^1$$
 R^2
 R^3
 R^4
 R^3
 R^4
 R^3
 R^4
 R^3
 R^5
 R^4
 R^3
 R^5

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an N-oxide form, a pharmaceutically acceptable addition salt, a quaternary amine and a stereochemically isomeric form thereof, wherein

- X represents O; $C(R^6)_2$ with R^6 being hydrogen, aryl or C_{1-6} alkyl optionally substituted with amino or mono- or di(C_{1-6} alkyl)amino; S or N- R^7 with R^7 being amino or hydroxy;
- R¹ represents C₁₋₆alkyl; aryl; thienyl; quinolinyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein the cycloC₃₋₁₂alkyl moiety optionally may contain a double bond and wherein one carbon atom in the cycloC₃₋₁₂alkyl moiety may be replaced by an oxygen atom or an NR⁸-moiety with R⁸ being hydrogen, benzyl or C₁₋₆alkyloxycarbonyl; wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyl, hydroxyC₁₋₆alkyl, haloC₁₋₆alkyl, aminoC₁₋₆alkyl, hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, halo, C₁₋₆alkyloxycarbonyl, aryl, amino, mono— or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, halo, piperazinyl, pyridinyl, morpholinyl, thienyl or a bivalent radical of formula –O-, -O-CH₂-O or –O-CH₂-CH₂-O-;

or a radical of formula (a-1)

$$Z_1$$
 CH Z_2 (CH₂)_n

a-1

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wherein Z₁ is a single covalent bond, O, NH or CH₂; Z₂ is a single covalent bond, O, NH or CH₂; n is an integer of 0, 1, 2 or 3;

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and wherein each hydrogen atom in the phenyl ring independently may optionally be replaced by halo, hydroxy, C_{1-6} alkyl, C_{1-6} alkyloxy or hydroxy C_{1-6} alkyl;

or X and R¹ may be taken together with the carbon atom to which X and R¹ are attached to form a radical of formula (b-1), (b-2) or (b-3);

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R² represents hydrogen; halo; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkylthio; C_{1-6} alkylcarbonyl; C_{1-6} alkyloxycarbonyl; C_{1-6} alkylcarbonyloxy C_{1-6} alkyl; 10 C₂₋₆alkenyl; hydroxyC₂₋₆alkenyl; C₂₋₆alkynyl; hydroxyC₂₋₆alkynyl; tri(C₁₋ 6alkyl)silaneC2-6alkynyl; amino; mono- or di(C1-6alkyl)amino; mono- or di(C₁₋₆alkyloxyC₁₋₆alkyl)amino; mono- or di(C₁₋₆alkylthioC₁₋₆alkyl)amino; aryl; arylC₁₋₆alkyl; arylC₂₋₆alkynyl; C₁₋₆alkyloxyC₁₋₆alkylaminoC₁₋₆alkyl; aminocarbonyl optionally substituted with C₁₋₆alkyl, C₁₋₆alkyloxyC₁₋₆alkyl, 15 C_{1-6} alkyloxycarbonyl C_{1-6} alkyl or pyridinyl C_{1-6} alkyl; a heterocycle selected from thienyl, furanyl, pyrrolyl, thiazolyl, oxazolyl, imidazolyl, isothiazolyl, isoxazolyl, pyrazolyl, pyridyl, pyrazinyl, pyridazinyl, pyrimidinyl, piperidinyl and piperazinyl, optionally N-substituted with C₁₋₆alkyloxyC₁₋₆alkyl, morpholinyl, thiomorpholinyl, dioxanyl or dithianyl; 20 a radical –NH-C(=O)R⁹ wherein R⁹ represents

C₁₋₆alkyl optionally substituted with cycloC₃₋₁₂alkyl, C₁₋₆alkyloxy, C₁₋₆alkyloxycarbonyl, aryl, aryloxy, thienyl, pyridinyl, mono- or di(C₁₋₆alkyl)amino, C₁₋₆alkylthio, benzylthio, pyridinylthio or pyrimidinylthio; cycloC₃₋₁₂alkyl; cyclohexenyl; amino; arylcycloC₃₋₁₂alkylamino; mono- or di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxycarbonylC₁₋₆alkyl)amino; mono- or di(c₁₋₆alkyloxycarbonyl)amino; mono- or di(arylC₁₋₆alkyl)amino; mono- or diarylamino; arylC₂₋₆alkenyl; furanylC₂₋₆alkenyl; piperididinyl; piperazinyl; indolyl; furyl; benzofuryl; tetrahydrofuryl; indenyl; adamantyl; pyridinyl; pyrazinyl; aryl; arylC₁₋₆alkylthio or a radical of formula (a-1);

a sulfonamid -NH-SO₂-R¹⁰ wherein R¹⁰ represents C_{1-6} alkyl, mono- or poly halo C_{1-6} alkyl, aryl C_{1-6} alkyl, aryl C_{2-6} alkenyl, aryl, quinolinyl, isoxazolyl or di $(C_{1-6}$ alkyl)amino;

- R³ and R⁴ each independently represent hydrogen; halo; hydroxy; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkyloxyC₁₋₆alkyl; C₁₋₆alkylcarbonyl; C₁₋₆alkyloxycarbonyl; C₂₋₆alkenyl; hydroxyC₂₋₆alkenyl; C₂₋₆alkynyl; hydroxyC₂₋₆alkynyl; tri(C₁₋₆alkyl)silaneC₂₋₆alkynyl; amino; mono– or di(C₁₋₆alkyl)amino; mono– or di(C₁₋₆alkyl)amino; aryl; morpholinylC₁₋₆alkyl or piperidinylC₁₋₆alkyl; or
 - R^2 and R^3 may be taken together to form $-R^2$ - R^3 -, which represents a bivalent radical of formula $-(CH_2)_3$ -, $-(CH_2)_4$ -, $-(CH_2)_5$ -, $-(CH_2)_6$ -, -CH=CH-CH=CH-, $-Z_4-CH=CH-$, $-CH=CH-Z_4$ -, $-Z_4-CH_2-CH_2-$, $-CH_2-CH_2-$, $-CH_2-CH_2-$, $-CH_2-CH_2-$,
- -CH₂-CH₂-CH₂-Z₄-, -Z₄-CH₂-CH₂-, -CH₂-Z₄-CH₂- or -CH₂-CH₂-Z₄-, with Z₄ being O, S, SO₂ or NR¹¹ wherein R¹¹ is hydrogen, C₁₋₆alkyl, benzyl or C₁₋₆alkyloxycarbonyl; and wherein each bivalent radical is optionally substituted with C₁₋₆alkyl.
 - or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH=CH-CH=CH- or -CH₂-CH₂-CH₂-CH₂-;
 - R⁵ represents hydrogen; cycloC₃₋₁₂alkyl; piperidinyl; oxo-thienyl; tetrahydrothienyl, arylC₁₋₆alkyl; C₁₋₆alkyloxyC₁₋₆alkyl; C₁₋₆alkyloxycarbonylC₁₋₆alkyl or C₁₋₆alkyl optionally substituted with a radical C(=O)NR_xR_y, in which R_x and R_y, each independently are hydrogen, cycloC₃₋₁₂alkyl, C₂₋₆alkynyl or C₁₋₆alkyl optionally substituted with cyano, C₁₋₆alkyloxy, C₁₋₆alkyloxycarbonyl, furanyl, pyrrolidinyl, benzylthio, pyridinyl, pyrrolyl or thienyl;
 - Y represents O or S;

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- or Y and R⁵ may be taken together to form =Y-R⁵- which represents a radical of formula
- 30 -CH=N-N= (c-1); -N=N-N= (c-2); or -N-CH=CH- (c-3);
 - aryl represents phenyl or naphthyl optionally substituted with one or more substituents selected from halo, hydroxy, C₁₋₆alkyl, C₁₋₆alkyloxy, phenyloxy, nitro, amino, thio, C₁₋₆alkylthio, haloC₁₋₆alkyl, polyhaloC₁₋₆alkyl, polyhaloC₁₋₆alkyloxy, hydroxyC₁₋₆alkyl, C₁₋₆alkyloxyC₁₋₆alkyl, aminoC₁₋₆alkyl, mono-or di(C₁₋₆alkyl)amino; mono-or di(C₁₋₆alkyl)aminoC₁₋₆alkyl, cyano, -CO-R¹², -CO-OR¹³,

-NR¹³SO₂R¹², -SO₂-NR¹³R¹⁴, -NR¹³C(O)R¹², -C(O)NR¹³R¹⁴, -SOR¹², -SO₂R¹²; wherein each R¹², R¹³ and R¹⁴ independently represent C₁₋₆alkyl; cycloC₃₋₆alkyl; phenyl; phenyl substituted with halo, hydroxy, C₁₋₆alkyl, C₁₋₆alkyloxy, haloC₁₋₆alkyl, polyhaloC₁₋₆alkyl, furanyl, thienyl, pyrrolyl, imidazolyl, thiazolyl or oxazolyl;

and when the R^1 -C(=X) moiety is linked to another position than the 7 or 8 position, then said 7 and 8 position may be substituted with R^{15} and R^{16} wherein either one or both of R^{15} and R^{16} represents C_{1-6} alkyl, C_{1-6} alkyloxy or R^{15} and R^{16} taken together may form a bivalent radical of formula -CH=CH-CH=CH-.

- 2. A compound according to claim 1, characterized in that,
- X represents O; $C(R^6)_2$ with R^6 being hydrogen or aryl; or N-R⁷ with R⁷ being amino or hydroxy;
- 15 R¹ represents C₁₋₆alkyl, aryl; thienyl; quinolinyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein the cycloC₃₋₁₂alkyl moiety optionally may contain a double bond and wherein one carbon atom in the cycloC₃₋₁₂alkyl moiety may be replaced by an oxygen atom or an NR⁸-moiety with R⁸ being benzyl or C₁₋₆alkyloxycarbonyl; wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyl, haloC₁₋₆alkyl, hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, halo, aryl, mono— or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, halo, piperazinyl, pyridinyl, morpholinyl, thienyl or a bivalent radical of formula —O- or -O-CH₂-CH₂-O-; or a radical of formula (a-1)

a-1

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wherein Z_1 is a single covalent bond, O or CH_2 ; Z_2 is a single covalent bond, O or CH_2 ; n is an integer of 0, 1, or 2; and wherein each hydrogen atom in the phenyl ring independently may optionally be replaced by halo or hydroxy;

or X and R¹ may be taken together with the carbon atom to which X and R¹ are attached to form a radical of formula (b-1), (b-2) or (b-3);

R² represents hydrogen; halo; cyano; C₁₋₆alkyl; C₁₋₆alkyloxy; C₁₋₆alkylthio; C₁₋₆alkylcarbonyl; C₁₋₆alkyloxycarbonyl; C₂₋₆alkenyl; hydroxyC₂₋₆alkenyl; C_{2-6} alkynyl; hydroxy C_{2-6} alkynyl; tri $(C_{1-6}$ alkyl)silane C_{2-6} alkynyl; amino; mono– or 5 di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxyC₁₋₆alkyl)amino; mono- or di(C₁₋₆alkylthioC₁₋₆alkyl)amino; aryl; arylC₁₋₆alkyl; arylC₂₋₆alkynyl; C_{1-6} alkyloxy C_{1-6} alkylamino C_{1-6} alkyl; aminocarbonyl optionally substituted with C₁₋₆alkyloxycarbonylC₁₋₆alkyl; a heterocycle selected from thienyl, furanyl, thiazolyl and piperidinyl, optionally 10 N-substituted with morpholinyl or thiomorpholinyl; a radical -NH-C(=0)R⁹ wherein R⁹ represents C₁₋₆alkyl optionally substituted with cycloC₃₋₁₂alkyl, C₁₋₆alkyloxy, C₁₋₆alkyloxycarbonyl, aryl, aryloxy, thienyl, pyridinyl, mono- or di(C₁₋₆alkyl)amino, C₁₋₆alkylthio, benzylthio, pyridinylthio or pyrimidinylthio; cycloC₃₋₁₂alkyl; cyclohexenyl; amino; arylcycloC₃₋₁₂alkylamino; 15 mono-or-di(C₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxycarbonylC₁₋₆alkyl)amino; mono- or di(C₁₋₆alkyloxycarbonyl)amino; mono-or di(C₂₋₆alkenyl)amino; mono- or di(arylC₁₋₆alkyl)amino; mono- or diarylamino; arylC₂₋₆alkenyl; furanylC₂₋₆alkenyl; piperididinyl; piperazinyl; indolyl; furyl; benzofuryl; tetrahydrofuryl; indenyl;

adamantyl; pyridinyl; pyrazinyl; aryl or a radical of formula (a-1); a sulfonamid -NH-SO $_2$ -R 10 wherein R 10 represents C $_{1\text{-}6}$ alkyl, mono- or poly haloC $_{1\text{-}6}$ alkyl, arylC $_{1\text{-}6}$ alkyl or aryl;

 R^3 and R^4 each independently represent hydrogen; C_{1-6} alkyl; C_{1-6} alkyloxy C_{1-6} alkyl; or

R² and R³ may be taken together to form –R²-R³-, which represents a bivalent radical of formula –(CH₂)₄-, –(CH₂)₅-, –Z₄-CH₂-CH₂-, –Z₄-CH₂-CH₂-CH₂- or -Z₄-CH₂-CH₂-, with Z₄ being O, S, SO₂ or NR¹¹ wherein R¹¹ is hydrogen, C₁₋₆alkyl, benzyl or C₁₋₆alkyloxycarbonyl; and wherein each bivalent radical is optionally substituted with C₁₋₆alkyl;

or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH=CH-CH=CH- or -CH₂-CH₂-CH₂-;

 R^5 represents hydrogen; piperidinyl; oxo-thienyl; tetrahydrothienyl, aryl C_{1-6} alkyl; C_{1-6} alkyloxycarbonyl C_{1-6} alkyl or C_{1-6} alkyl optionally substituted with a radical

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C(=O)NR_xR_y, in which R_x and R_y, each independently are hydrogen, cycloC₃₋₁₂alkyl, C₂₋₆alkynyl or C₁₋₆alkyl optionally substituted with cyano, C₁₋₆alkyloxy or C₁₋₆alkyloxycarbonyl;

Y represents O or S;

or Y and R⁵ may be taken together to form =Y-R⁵- which represents a radical of formula

-CH=N-N=

(c-1); or

-N=N-N=

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(c-2);

aryl represents phenyl or naphthyl optionally substituted with one or more substituents selected from halo, C₁₋₆alkyloxy, phenyloxy, mono-or di(C₁₋₆alkyl)amino and cyano;

and when the R^1 -C(=X) moiety is linked to another position than the 7 or 8 position, then said 7 and 8 position may be substituted with R^{15} and R^{16} wherein either one or both of R^{15} and R^{16} represents C_{1-6} alkyl or R^{15} and R^{16} taken together may form a bivalent radical of formula -CH=CH-CH=CH-.

- 3. A compound according to claim 1, characterized in that,
- X represents O;
- R¹ represents C₁₋₆alkyl; cycloC₃₋₁₂alkyl or (cycloC₃₋₁₂alkyl)C₁₋₆alkyl, wherein one or more hydrogen atoms in a C₁₋₆alkyl-moiety or in a cycloC₃₋₁₂alkyl-moiety optionally may be replaced by C₁₋₆alkyloxy, aryl, halo or thienyl;
- R² represents hydrogen; halo; C₁₋₆alkyl or amino;
- R³ and R⁴ each independently represent hydrogen or C₁₋₆alkyl; or
- R² and R³ may be taken together to form –R²-R³-, which represents a bivalent radical of formula –Z₄-CH₂-CH₂- or -Z₄-CH₂-CH₂- with Z₄ being O or NR¹¹ wherein R¹¹ is C₁₋₆alkyl; and wherein each bivalent radical is optionally substituted with C₁₋₆alkyl;

or R³ and R⁴ may be taken together to form a bivalent radical of formula -CH₂-CH₂-CH₂-CH₂-;

30 R⁵ represents hydrogen;

Y represents O; and

aryl represents phenyl optionally substituted with halo.

- 4. A compound as claimed in claim 1, characterized in that, the R¹-C(=X) moiety is linked to the quinoline or quinolinone moiety in position 6.
 - 5. A compound as claimed in claim 1 for use as a medicine.

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- 6. Use of a compound as defined in claims 1 to 4 in the manufacture of a medicament for treating or preventing glutamate-induced diseases of the central nervous system.
- Use according to claim 6, characterized in that, the glutamate-induced disease of the central nervous system is drug addiction or abstinence (dependence, opioid tolerance, opioid withdrawal), hypoxic, anoxic and ischemic injuries (ischemic stroke, cardiac arrest), pain (neuropathic pain, inflammatory pain, hyperalgesia), hypoglycemia, diseases related to neuronal damage, brain trauma, head trauma, spinal cord injury, myelopathy, dementia, anxiety, schizophrenia, depression, impaired cognition, amnesia, bipolar disorders, conduct disorders, Alzheimer's disease, vascular dementia, mixed (Alzheimer's and vascular) dementia, Lewy Body disease, delirium or confusion, Parkinson's disease, Huntington's disease, Down syndrome, epilepsy, aging, Amyotrophic Lateral Sclerosis, multiple sclerosis, AIDS (Acquired Immune Deficiency Syndrome) and AIDS related complex (ARC).
 - 8. A pharmaceutical composition comprising a pharmaceutically acceptable carrier, and as active ingredient a therapeutically effective amount of a compound as defined in claims 1 to 4.
 - 9. A process of preparing a composition as claimed in claim 8, characterized in that, a pharmaceutically acceptable carrier is intimately mixed with a therapeutically effective amount of a compound as described in claims 1 to 4.
- 25 10. A process of preparing a compound of formula (I-A) or (I-B) as claimed in claim 1, characterized by
 - a) oxidizing an intermediate of formula (II) in the presence of a suitable oxidizing agent

$$R^{1}$$
— CH — Q oxidation R^{1} — C — Q

(II) $(I_{A/B}$ - $a)$

with R¹ as defined in claim 1 and Q representing the quinoline or the quinoline moiety of a compound of formula (I-A) or (I-B); or

b) reacting an intermediate of formula (III) with an intermediate of formula (IV)

$$R^1$$
— C = N + W_1 — Q \longrightarrow R^1 — C — Q (III) (IV) (I_{A/B}-a)

with R^1 as defined in claim 1, Q representing the quinoline or the quinoline moiety of a compound of formula (I-A) or (I-B) and W_1 being a suitable leaving group; or c) reacting an intermediate of formula (V) with an intermediate of formula (IV)

$$R^{1} = C - N$$
 CH_{3}
 (IV)
 $R^{1} = C - Q$
 $(I_{A/B}-a)$

with R¹ as defined in claim 1, Q representing the quinoline or the quinolinone moiety of a compound of formula (I-A) or (I-B) and W₁ being a suitable leaving group; or d) reacting an intermediate of formula (VI) with an intermediate of formula (VII) in the presence of a suitable acid

$$R^{1a}$$
 OH + HO C Q \rightarrow R^{1a} O C Q (VI) (VII)

with R^{1a} being defined as R¹ according to claim 1 provided that R¹ is linked to the carbonyl moiety via a oxygen atom and Q representing the quinoline or the quinolinene moiety of a compound of formula (I-A) or (I-B); or

e) reacting an intermediate of formula (VIII) in the presence of a suitable acid

with R^1 , X, R^3 and R^4 defined as in claim 1;

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and, if desired, converting compounds of formula (I-A) or (I-B) into each other following art-known transformations; and further, if desired, converting the compounds of formula (I-A) or (I-B), into a therapeutically active non-toxic acid addition salt by treatment with an acid, or conversely, converting the acid addition salt form into the free base by treatment with alkali; and, if desired, preparing stereochemically isomeric forms, quaternary amines or *N*-oxide forms thereof.

INTERNATIONAL SEARCH REPORT Interional Application No PCT/EP 01/11135 A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07D215/18 C07D215/22 CO7D215/14 C07D215/38 C07F7/10 CO7D215/48 CO7D409/04 CO7D471/04 CO7D215/36 C07D405/04 CO7D491/04 CO7D495/04 A61K31/47 CO7D409/12 CO7D401/12 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 CO7D CO7F A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ° Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. US 4 476 132 A (GOESCHKE RICHARD 1,2,4-6,χ 9 October 1984 (1984-10-09) 8,9 see general formula and examples WO 97 44339 A (GOULET MARK ; JIANG JINLONG 1,2,4 χ (US); ALLEN ERIC E (US); DEVITA ROBERT) 27 November 1997 (1997-11-27) see formula 5, page 13 and defintions of

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X Further documents are listed in the continuation of box C.	χ Patent family members are listed in annex.
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Date of the actual completion of the international search 26 November 2001	Date of mailing of the international search report 30/11/2001
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswljk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Scruton-Evans, I

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